Remarks

Reconsideration of this Application is respectfully requested.

Upon entry of the foregoing amendment, claims 1, 5-13, 18 and 20-29 are pending in the application, with claim 1 being the independent claim. Claims 2-4, 14-17, and 19 are sought to be cancelled without prejudice to or disclaimer of the subject matter therein. New claims 20-29 are sought to be added. Claim 1 has been amended to be directed to Applicants' elected peptides and to clarify the claimed invention. New claims 20-29 depend from claim 1 and are each directed to one of Applicants' elected peptides. Support for claim 1 and new claims 20-29 can be found, *inter alia*, at ¶ 41 and Tables 11-18. Claims 5-9 have been amended to clarify the claimed invention. Support for amended claims 5 and 6 can be found, *inter alia*, at ¶ 69. Support for amended claims 7 and 9 can be found, *inter alia*, at ¶ 98. Support for amended claim 8 can be found, *inter alia*, at ¶ 73. Claim 13 has been amended to be directed to Applicants' elected antigen. These changes are believed to introduce no new matter, and their entry is respectfully requested.

Based on the above amendment and the following remarks, Applicants respectfully request that the Examiner reconsider all outstanding objections and rejections and that they be withdrawn.

Objection to the Oath/Declaration

The Examiner has objected to the Oath/Declaration of April 5, 2005, citing that non-initialed and/or non-dated alterations have been made to the oath or declaration. (Office Action, ¶ 6, Page 3.) Applicants note that the Declaration submitted to the U.S.

Receiving Office upon filing of the corresponding International Application did not contain any such alterations. Applicants are resubmitting herewith a copy of the Declaration as originally submitted.

It appears that the alterations in the scanned version of the Declaration on the PAIR system were made to the document by USPTO personnel subsequent to the submission of the Declaration by Applicants. Applications respectfully request that the Examiner consider the resubmitted Declaration, which does not contain any alterations, and withdraw the outstanding objection.

Objection to the Abstract

The Examiner has objected to the abstract of the disclosure. (Office Action, ¶ 8, Page 3.) Applicants have amended herewith the abstract of the disclosure. Accordingly, Applicants respectfully request that the Examiner reconsider and withdraw the outstanding objection.

Objections to the Specification

The Examiner has objected to the specification because it does not provide sequence identifiers for the peptide sequences of ≥ four (4) amino acid residues in length pursuant to 37 C.F.R. § 1.821 (c) and/or (d), for example, in Tables 5, 6, and 11-29. (Office Action, ¶ 9, Page 3.)

Applicants have submitted herewith replacement pages for Tables 11-29 that include sequence identifiers as required by 37 C.F.R. §§ 1.821 (c) and/or (d).

Applicants have also submitted herewith replacement pages for Tables 5 and 6. Applicants note that the amino acids listed in Tables 5 and 6 refer to amino acids that can be present in the alternative at certain anchor residue positions (p1=position 1; p4=position 4; and p6=position 6) of an HLA-DR core motif for the HLA alleles DR supertype, DR3a or DR3b. The amino acids are referred to by their standard single letter designations. As an example, for the DR3 supertype, the anchor residues at position (p1) can be, in the alternative, a leucine (L), an isoleucine (I), a valine (V), a methionine (M), a phenylalanine (F), a tryptophan (W), or a tyrosine(Y). All of the amino acids listed in Tables 5 and 6 are alternative amino acids at the positions indicated according to the designation described above.

In order to clarify that these amino acids are present in the alternative at the positions indicated, Applicants have amended Tables 5 and 6 to include a comma between each amino acid. Because the amino acids in Tables 5 and 6 correspond to individual amino acids, they are not peptide sequences of \geq 4 amino acid residues in length, and thus no corresponding sequence identifier is required pursuant to 37 C.F.R. §§ 1.821 (e) and/or (d).

The Examiner has further objected to the legend of Figure 1 because it does not recite the sequence identifiers for the anchor residues having \geq four (4) amino acid residues shown in Figure 1. (Office Action, ¶ 10, Page 3.) Applicants note that the amino acids shown in Figure 1, similar to those shown in Tables 5 and 6, refer to amino acids that can be present in the alternative at the designated anchor residue positions, and thus refer to single amino acids, rather than a peptide sequence having \geq four (4) amino

acid residues. Applicants have amended the figure legend to Figure 1 herewith to clarify this.

Accordingly, based on the above amendments to the specification, Applicants respectfully request that the Examiner reconsider and withdraw the outstanding objections.

Rejections under 35 U.S.C. § 112, second paragraph

Claims 1, 3-15 and 18 have been rejected under 35 U.S.C. § 112, second paragraph, as allegedly being indefinite for failing to particularly point out and distinctly claim the subject matter which Applicants regard as their invention. (Office Action, ¶ 11, Page 3.)

The Examiner has alleged that claims 1, 3-15 and 18 are indefinite for reciting "peptides" because in claim 1 it is not clear what the metes and bounds of the size of the peptide is. (Office Action, ¶ 11a, Page 3.) Without acquiescing to the Examiner's rejection and solely in efforts to expedite prosecution, Applicants have amended claim 1 to clarify the claimed invention. Applicants assert that the claims as amended are clear as to the invention's metes and bounds.

The Examiner has also alleged that in claims 3 and 5, the phrase "HTL epitope" is not defined by the claims or in the specification, and it is not clear what the metes and bounds of the phrase are with respect to the overall composition. (Office Action, ¶ 11b, Page 3.) Without acquiescing to the Examiner's rejection and solely in efforts to expedite prosecution, Applicants note that claim 3 has been cancelled. Applicants also assert that the phrase "HTL epitope" is clear from the specification. The specification

discloses that HTL refers to "helper T lymphocyte" and describes peptide fragments that activate such helper T lymphocytes:

> MHC molecules are classified as either class I or class II. Class II MHC molecules are expressed primarily on activated lymphocytes and antigen-presenting cells. CD4+ T lymphocytes are activated with recognition of a unique peptide fragment presented by a class II MHC molecule, usually found on an antigen presenting cell like a macrophage or dendritic cell. Often known as helper T lymphocytes (HTL), CD4+ lymphocytes proliferate and secrete cytokines that either support a antibody-mediated response through the production of IL-4 and IL-10 or support a cell-mediated response through the production of IL-2 and IFN-v.

(Specification, ¶ 2 (emphasis added).) The specification further describes that the unique peptide fragment presented by a class II MHC molecule is an "epitope" of the native antigen:

> The MHC-binding peptides identified herein represent epitopes of a native antigen. With regard to a particular amino acid sequence, an epitope is a set of amino acid residues which is recognized by a particular antibody or T cell receptor. Such epitopes are usually presented to lymphocytes via the MHC-peptide complex. An epitope retains the collective features of a molecule, such as primary, secondary and tertiary peptide structure, and charge, that together form a site recognized by an antibody, T cell receptor or MHC molecule.

(Specification, ¶ 35.) In addition, the specification describes that "[olne embodiment of an HTL-inducing peptide is less than 50 residues in length and usually consist of between about 6 and about 30 residues, more usually between about 12 and 25, and often between about 15 and 20 residues, for example 15, 16, 17, 18, 19, or 20 residues." (Specification, ¶ 41.) Specific examples, of such HTL-inducing peptides are disclosed, for example, in Tables 23-27. Thus, as shown above, the specification clearly sets forth

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what is meant by the phrase "HTL epitope" and discloses specific examples of such HTL epitopes. Without acquiescing to the Examiners rejection and solely in efforts to expedite prosecution, Applicants have amended claim 5 to clarify the metes and bounds of the phrase with respect to the overall composition.

The Examiner has further alleged that claims 3-5 and 8 are indefinite because it is not clear how the HTL and CTL epitope and the MHC targeting sequences are related to the peptides of claim 1. (Office Action, ¶ 11c, Page 3.) Without acquiescing to the Examiner's rejection, Applicants have cancelled claims 3 and 4. Applicants note that the limitation of claim 4 has been incorporated into amended claim 1. Applicants have amended claims 5 and 8 to clarify the relationship of the HTL epitope and the MHC targeting sequence to the peptides of claim 1.

The Examiner has further alleged that claims 1, 6, 7 and 9 are indefinite as to how the "spacer molecule" (Claim 6), "carrier: (Claim 7) and "lipid" (Claim 9) relate to the peptides of the composition of claim 1. (Office Action, ¶ 11d, Page 3.) Without acquiescing to the Examiner's rejection, Applicants have amended claims 6, 7, and 9 to clarify the relationship of the spacer molecule, carrier and lipid to the peptides of claim 1.

The Examiner has also alleged that claims 11 and 12 are indefinite because the terms "heteropolymer" and "homopolymer" are not defined in the specification, and it is not clear what the metes and bounds of the terms are with respect to the overall composition. (Office Action, ¶ 11e, Page 3.) The specification states that peptide(s) of the invention may be:

... linked to its own carrier or as a homopolymer or heteropolymer of active peptide units. Such a polymer has the advantage of increased immunological reaction and, where different peptides are used to make up the polymer, the additional ability to induce antibodies and/or CTLs that react with different antigenic determinants of the virus or tumor cells.

(Specification, ¶ 98.) Thus, the specification discusses that the units of a homopolymer or heteropolymer are the peptides of the invention. Furthermore, the terms "homo" and "hetero" are clear on their face, and would be clearly understand by one of ordinary skill in the art to mean "same" and "different," respectively. Thus, one of ordinary skill in the art would understand that a "homopolymer" is made up of the same polymer units, and a "heteropolymer" is made up of different polymer units. In view of the specification describing that the units of such polymers are individual peptides of the invention, it can be readily understood from the specification that the meaning of "homopolymer" is a polymer with each unit composed of the same peptide, whereas a "heteropolymer" is a polymer with units composed of different peptides.

In view of the discussion above, Applicants respectfully assert that the pending claims comply with the requirements of 35 U.S.C. § 112, second paragraph. Accordingly, Applicants respectfully request that the Examiner reconsider and withdraw the outstanding rejections.

Rejections under 35 U.S.C. § 112, first paragraph

Claims 3-5, 8, 11 and 12 are rejected under 35 U.S.C. § 112, first paragraph, as allegedly failing to comply with the enablement requirement. (Office Action, ¶ 12, Page 6.) Applicants respectfully disagree and traverse the rejection.

As an initial matter, without acquiescing to the Examiner's rejection and solely in efforts to expedite prosecution, Applicants have cancelled claims 3 and 4. As discussed further below. Applicants assert that claims 5. 8. 11 and 12 are enabled.

The Examiner has alleged that

The specification does not show that any of the elected peptides has any of the claimed functional properties, namely, CTL-induction, HTL-induction or MHC targeting. The specification does not provide a single working example for any one of the elected peptides demonstrating that the peptide has one or more of the claimed functional properties in a relevant bioassay or animal model.

(Office Action, Page 7, ¶ 12.)

"As concerns the breadth of a claim relevant to enablement, the only relevant concern should be whether the scope of enablement provided to one skilled in the art by the disclosure is commensurate with the scope of protection sought by the claims." MPEP § 2164.08 (2006) (citing AK Steel Corp. v. Sollac, 344 F.3d 1234, 1244 (Fed. Cir. 2003); In re Moore, 439 F.2d 1232, 1236 (C.C.P.A. 1971); see also Plant Genetic Sys., N.V. v. DeKalb Genetics Corp., 315 F.3d 1335, 1339 (Fed. Cir. 2003).

Applicants note that the pending claims, as indicated by the Examiner, are directed to a *composition* comprising one or more peptides selected from the peptides of SEQ ID NOs: 53, 55, 139, 502, 527, 627, 673, 807, 846 and 859. (Office Action, ¶ 12, Page 6, "Nature of the Invention.") Applicants point out that the claims do not require any further limitations. Although the Examiner refers to the "claimed" functional properties of the elected peptides, Applicants again note that the claims are only directed to a composition comprising one or more of the elected peptides and do not recite any further claimed features.

Moreover, as long as the specification discloses at least one method for making and using the claimed invention that bears a reasonable correlation to the entire scope of the claim, then the enablement requirement of 35 U.S.C. § 112 is satisfied. *In re Fisher*, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970). Additionally, "a specification disclosure which contains a teaching of the manner and process of making and using the invention must be taken as in compliance with the enabling requirement of the first paragraph of § 112 unless there is reason to doubt the objective truth of the statements contained therein which must be relied upon for enabling support." *Rasmusson v. Smithkline Beecham Corp.*, 413 F.3d 1318, 1323 (Fed. Cir. 2005) (quoting *In re Marzocchi*, 439 F.2d 220, 223 (C.C.P.A. 1971)).

Applicants assert that the specification discloses how to make and use compositions comprising the elected peptides and how to test these compositions for binding affinity using various assays. (Specification, ¶ 55-59.) The specification also discloses how to test these compositions for inhibition of CTL or HTL recognition using purified MHC molecules and radioiodonated peptides and/or cells by immunofluorescent staining and flow microfluorometry, or peptide-dependent class I assembly assays. (Specification, ¶ 77.) The specification also discusses the use of mutant mammalian cell lines to test for the capacity of compositions comprising a particular peptide to induce *in vitro* primary CTL responses. (Specification, ¶ 78.) Additionally, the specification discloses how such compositions can be made. (See Specification, ¶ 61-75.)

Thus, as discussed above, the specification not only discloses how to make and use compositions comprising Applicants' elected peptides, the specification also teaches how to assay the efficacy of such compositions and determine whether the particular

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peptides have the capacity to bind to certain MHC molecules and/or have the ability to induce an immune response. Thus, the specification discloses "at least one method for making and using the claimed invention that bears a reasonable correlation to the entire scope of the claim."

The purpose of the requirement that the specification describe the invention in such terms that one skilled in the art can make and use the claimed invention without undue experimentation (i.e., the enablement requirement) is to ensure that the invention is communicated to the interested public in a meaningful way. Here, Applicants respectfully assert that the information contained in the disclosure of the specification is sufficient to inform those skilled in the relevant art how to both make and use the claimed invention in a manner that satisfies the requirements of 35 U.S.C. § 112, first paragraph for enablement.

Finally, the Examiner cites several documents referring to the "inherent risks associated with T-cell immunogenic peptides in general." (Office Action, ¶ 12, Pages 7-9.) The fact that experimentation may be complex does not necessarily make it undue, if the art typically engages in such experimentation. In re Certain Limited-Charge Cell Culture Microcarriers, 221 USPQ 1165, 1174 (Int'l Trade Comm"n 1983), aff'd. sub nom., Massachusetts Institute of Technology v. A.B. Fortia, 774 F.2d 1104, 227 USPQ 428 (Fed. Cir. 1985). See also In re Wands, 858 F.2d at 737, 8 USPQ2d at 1404. Here Applicants respectfully assert that not only is no experimentation necessary to practice the claimed invention, but even if any experimentation was performed it would not be undue as it would be of the type typically engaged in by artisans in this art.

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Claims 14 and 15 are rejected under 35 U.S.C. § 112, first paragraph, as allegedly failing to comply with the enablement requirement. (Office Action, ¶13, Page 6.) Without acquiescing to the Examiner's rejection and solely in efforts to expedite prosecution, Applicants have cancelled claims 14 and 15. The rejections of these claims has therefore been rendered moot. Accordingly, Applicants respectfully request that the Examiner reconsider and withdraw the outstanding rejection.

Other Matters

Applicants note that the Examiner has indicated that Applicants' elected peptides (SEQ ID NOs: 53, 55, 139, 502, 527, 627, 673, 807, 846 and 859) are free of the prior art.

Conclusion

All of the stated grounds of objection and rejection have been properly traversed, accommodated, or rendered moot. Applicants therefore respectfully request that the Examiner reconsider all presently outstanding objections and rejections and that they be withdrawn. Applicants believe that a full and complete reply has been made to the outstanding Office Action and, as such, the present application is in condition for allowance. If the Examiner believes, for any reason, that personal communication will expedite prosecution of this application, the Examiner is invited to telephone the undersigned at the number provided.

Prompt and favorable consideration of this Amendment and Reply is respectfully requested.

Respectfully submitted,

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usually consists of about 8, 9, 10 or 11 residues, preferably 9 or 10 residues. In one embodiment, HLA-DR3 a binding is characterized by an L, I, V, M, F or Y residue at position 1 and a D or E residue at position 4. In another embodiment, HLA-DR3 b binding is characterized by an L, I, V, M, F, Y or A residue at position 1, a D, E, N, Q, S or T residue at position 4, and a K, R or H residue at position 6. In another embodiment, key anchor residues of a DR supertype binding motif are an L, I, V, M, F, W or Y residue at position 1 and an L, I, V, M, S, T, P, C or A residue at position 6. See table 5.

TABLE 5
HLA-DR motifs

	Anchor residues of HLA-DR core motifs						
•	p1	p4	р6				
DR supertype	LIVMFWY		LIVMSTPCA				
	L,I,V,M,F,W,Y		L,I,V,M,S,T,P,C,A				
DR3 a	LIVMFY	ĐE					
	L,I,V,M,F,Y	D,E					
DR3 b	LIVMFYA	DENQST	KRH				
	L,I,V,M,F,Y,A	D,E,N,Q,S,T	<u>K,R,H</u>				

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Moreover, in another embodiment, murine Db binding is characterized by an N residue at position 5 and L, I, V or M residue at the C-terminal position. In yet another embodiment, murine Kb binding is characterized by a Y or F residue at position 5 and an L, I, V or M residue at the C-terminal position. In an additional embodiment, murine Kd binding is characterized a Y or F residue at position 2 and an L, I, V, or M residue at the C-terminal position. In a further embodiment, murine Kk binding is characterized by an E or D residue at position 2 and an L, I, M, V, F, W, Y or A residue at the C-terminal position. In a further embodiment, murine Ld binding is characterized by a P residue at position 2 and an L, I, M, V, F, W or Y residue at the C-terminal position. See Table 6.

Table 6
Murine Class I Motifs

	Anchor residues of mouse class I motifs				
Allele	p2	р3	р5	C terminus	
 Db		-	N	LIVM <u>L,I,V,M</u>	
 Dd	G	P		LVI <u>L,V,I</u>	
Kb			YF	LVI ML,V,I,M	
Kd	YF			LVI ML,V,I,M	
Kk	ED			LIMVA <u>L,I,M,V,A</u>	
Ld	P			LIMVFWYL,I,M,V,F,W,	

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The peptides present in the invention can be identified by any suitable method. For example, peptides are conveniently identified using the algorithms of the invention described in the co-pending U.S. Patent Application Serial No. 09/894,018. These algorithms are mathematical procedures that produce a score which enables the selection of immunogenic peptides. Typically one uses the algorithmic score with a binding threshold to enable selection of peptides that have a high probability of binding at a certain affinity and will in turn be immunogenic. The algorithm are based upon either the effects on MHC binding of a particular amino acid at a particular position of a peptide or the effects on binding MHC of a particular substitution in a motif containing peptide.

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Peptide sequences characterized in molecular binding assays and capture assays have been and can be identified utilizing various technologies. Motif-positive sequences are identified using a customized application created at Epimmune. Sequences are also identified utilizing matrix-based algorithms, and have been used in conjunction with a "power" module that generates a predicted 50% inhibitory concentration (PIC) value. These latter methods are operational on Epimmune's HTML-based Epitope Information System (EIS) database. All of the described methods are viable options in peptide sequence selection for ICso determination using binding assays.

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Additional procedures useful in identifying the peptides of the present invention generally follow the methods disclosed in Falk et al., Nature

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TABLE 11

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Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
AYGPGPGKF	1	9	Artificial sequence			A
AEIPYLAKY	2	9	Artificial sequence	pool consensus		Α
AADAAAAKY	<u>3</u>	9	Artificial sequence			PolyA
AYSSWMYSY	4	9	EBV	EBNA3	176	
LAEKTMKEY	<u>5</u>	9	FluA	POL2	16	
GTYDYWAGY	<u>6</u>	9	Gonorrhea			
LSVHSIQNDY	7	10	Gonorrhea			
DTGQCPELVY	8	10	Gonorrhea			
DLLDTASALY	2	10	HBV	Core	419	
WFHISCLTF	10	9	HBV	NUC	102	
LSLDVSAAFY	11	10	HBV	pol	426	
LSGPGPGAFY	12	10	HBV	pol	426	Α
LSLGPGPGFY	13	10	HBV	pol	426	Α
LSLDGPGPGY	14	10	HBV	pol	426	Α
KTYGRKLHLY	15	10	HBV	pol	1098	
KTGPGPGHLY	16	10	HBV	pol	1098	Α
KTYGPGPGLY	17	10	HBV	pol	1098	Α
KTYGGPGPGY	18	10	HBV	pol	1098	Α
KYTSFPWL	19	8	HBV	pol	745	
FAAPFTQCGY	20	10	HBV	pol	631	
SYOHFRKLLL	21	10	HBV	POL	4	
LYSHPIILGF	22	10	HBV	POL	492	
MSTTDLEAY	23	9	HBV	x	103	
MYVGGPGPGVF	24	11	HCV	El	275	Α
VMGSSYGF	25	8	HCV	NS5	2639	
EVDGVRLHRY	26	10	HCV	NS5	2129	
RTEILDLWVY	27	10	HIV	NEF	182	Α
RQDILDLWVY	28	10	HIV	NEF	182	Α
RTDILDLWVY	29	10	HIV	NEF	182	Α
YTDGPGIRY	30	9	HIV	NEF	207	Α
ATELHPEYY	31	9	HIV	NEF	322	Α
DLWVYHTQGYY	32	11	HIV	NEF	188	Α
WVYHTQGYY	33	9	HIV	NEF	191	A
FFLKEKGGF	34	9	HIV	NEF	116	Α
LYVYHTQGY	35	9	HIV	NEF	190	A
ITKILYQSNPY	3.6	11	HIV	REV	20	Α
KTLYQSNPY	37	9	HIV	REV	22	Α
PVDPNLEPY	38	9	HIV	TAT	3	A
STVKHHMY	39	8	HIV	VIF	23	Α
LSKISEYRHY	40	10	HPV	E6	70	
ISEYRHYNY	41	9	HPV	E6	73	
RFHNIRGRW	42	9	HPV	E6	131	
RFLSKISEY	43	9	HPV	E6	68	

HLA-A1 SUPERTYPE

-	SEQ ID					
Sequence	NO.	AA	Organism	Protein		Analog
TLEKLTNTGLY	<u>45</u>	11	HPV	E6	89	
TLGPGPGTGLY	<u>46</u>	11	HPV	E6		A
TLEGPGPGGLY	<u>47</u>	11	HPV	E6		A
TLEKGPGPGLY	<u>48</u>	11	HPV	E6	89	A
TLEKLGPGPGY	<u>49</u>	11	HPV	E6	89	A
TLEKLTNTGLY	<u>50</u>	11	HPV	E6	89	
TLEKITNTELY	<u>51</u>	11	HPV	E6	89	
PYGVCIMCLRF	<u>52</u>	11	HPV	E6	59	
ITDIILECVY	<u>53</u>	10	HPV	E6	30	Α
YSDISEYRHY	<u>54</u>	10	HPV	E6	77	A
LTDIEITCVY	<u>55</u>	10	HPV	E6	25	A
YSDIRELRHY	<u>56</u>	10	HPV	E6	72	Α
ELSSALEIPY	<u>57</u>	10	HPV	E6	14	
ETSSALEIPY	58	10	HPV	E6	14	A
ELDSALEIPY	59	10	HPV	E6	14	A
YTKVSEFRWY	60	10	HPV	E6	70	A
YSDVSEFRWY	<u>61</u>	10	HPV	E6	70	A
LTDVSIACVY	<u>62</u>	10	HPV	E6	25	A
FTSRIRELRY	<u>63</u>	10	HPV	E6	71	A
YSDIRELRYY	64	10	HPV	E6	72	A
LTDLRLSCVY	<u>65</u>	10	HPV	E6	26	A
FTSKVRKYRY	<u>66</u>	10	HPV	E6	72	A
YSDVRKYRYY	<u>67</u>	10	HPV	E6	73	A
FYSKVSEFRF	<u>68</u>	10	HPV	E6	69	A
FYSRIRELRF	<u>69</u>	10	HPV	E6	71	A
PYAVCRVCLF	<u>70</u>	10	HPV	E6	62	Α
ITEYRHYNY	<u>71</u>	9	HPV	E6	73	A
ISDYRHYNY	<u>72</u>	9	HPV	E6	73	A
ITEYRHYQY	<u>73</u>	9	HPV	E6	73	A
ISDYRHYQY	<u>74</u>	9	HPV	E6	73	A
LTDLLIRCY	<u>75</u>	9	HPV	E6	99	A
KTDQRSEVY	<u>76</u>	9	HPV	E6	35	A
AYRDLCIVY	<u>77</u>	9	HPV	E6	53	A
KYYSKISEY	<u>78</u>	9	HPV	E6	75	A
KFYSKISEF	<u>79</u>	9	HPV	E6	75	A
RYHNIRGRW	80	9	HPV	E6	131	A
RFHNIRGRF	<u>81</u>	9	HPV	E6	131	A
AYKDLFVVY	<u>82</u>	9	HPV	E6	48	Α
LFVVYRDSF	<u>83</u>	9	HPV	E6	52	A
RYHNIAGHY	<u>84</u>	9	HPV	E6	126	A
RFHNIAGHF	<u>85</u>	9	HPV	E6	126	A
VYGTTLEKF	<u>86</u>	9	HPV	E6	83	A
AYADLTVVY	<u>87</u>	9	HPV	E6	46	A
AFADLTVVF	88	9	HPV	E6	46	A
RYLSKISEY	<u>89</u>	9	HPV	E6	68	A
RYHNISGRW	<u>90</u>	9	HPV	E6	124	A

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Sequence	SEQ ID NO.	AA	Organism	Protein	Position Analog
AYKDLCIVY	91	9	HPV	E6	48 A
RYHSIAGQY	92	9	HPV	E6	126 A
RFHSIAGQF	<u>93</u>	9	HPV	E6	126 A
KYLFTDLRI	<u>94</u>	9	HPV	E6	44 A
KFLFTDLRF	95	9	HPV	E6	44 A
LYTDLRIVY	<u>96</u>	9	HPV	E6	46 A
LFTDLRIVF	<u>97</u>	9	HPV	E6	46 A
RFLSKISEF	<u>98</u>	9	HPV	E6	68 A
EYRHYQYSF	<u>99</u>	9	HPV	E6	75 A
RYHNIMGRW	100	9	HPV	E6	124 A
RFHNIMGRF	101	9	HPV	E6	124 A
NFACTELKF	102	9	HPV	E6	47 A
PYAVCRVCF	103	9	HPV	E6	62 A
LYYSKVRKY	104	9	HPV	E6	71 A
VYADLRIVY	105	9	HPV	E6	46 A
VFADLRIVF	106	9	HPV	E6	46 A
NYSLYGDTF	107	9	HPV	E6	80 A
RFHNISGRF	108	9	HPV	E6	124 A
FTDLTIVY	109	8	HPV	E6	47
FTDLRIVY	110	8	HPV	E6	47
TLEKLTNTGLY	111	11	HPV	E6	89
LTDIEITCVY	112	10	HPV	E6	25 A
LTDVSIACVY	113	10	HPV	E6	25 A
ITDIILECVY	114	10	HPV	E6	30
KTDQRSEVY	115	9	HPV	E6	35
FTDLTIVY	116	8	HPV	E6	47
YSDIRELRYY	117	10	HPV	E6	72 A
YTKVSEFRWY	118	10	HPV	E6	70 A
FTSRIRELRY	119	10	HPV	E6	71 A
FTSKVRKYRY	120	10	HPV	E6	72 A
ISDYRHYNY	121	9	HPV	E6	73 A
ISEYRHYQY	122	9	HPV	E6	73
ISDYRHYQY	123	9	HPV	E6	73 A
EYRHYCYSLY	124	10	HPV	E6	82
EYRHYNYSLY	125	10	HPV	E6	75
LTDLLIRCY	126	9	HPV	E6	99
ETRHYCYSLY	127	10	HPV	E6	82 A
EYDHYCYSLY	128	10	HPV	E6	82 A
KTRYYDYSVY	129	10	HPV	E6	78 A
KYDYYDYSVY	<u>130</u>	10	HPV	E6	78 A
ETRHYNYSLY	131	10	HPV	E6	75 A
EYDHYNYSLY	<u>132</u>	10	HPV	E6	75 A
PTLKEYVLDLY	133	11	HPV	E7	6
HTDTPTLHEY	<u>134</u>	10	HPV	E7	2 A
RTETPTLQDY	<u>135</u>	10	HPV	E7	2 A
ETDPVDLLCY	136	10	HPV	E7	20 A

HLA-A1 SUPERTYPE

_	SEQ ID			B	D. states	4
Sequence QTEQATSNYY	NO. 137	10	Organism HPV	Protein E7	Position 46	Analog A
ATDNYYIVTY	138	10	HPV	E7	50	A
	138	9	HPV	E7 E7	8	A
LTEYVLDLY		9		E7	46	A
QTEQATSNY	<u>140</u>		HPV			А
RQAKQHTCY	<u>141</u>	9	HPV	E7	51	
RTAKQHTCY	142	9	HPV	E7	51	A
HTDTPTLHEY	143	10	HPV	E7	2	A
RTETPTLQDY	144	10	HPV	E7	2	Α
PTLKEYVLDLY	145	11	HPV	E7	6	
LTEYVLDLY	<u>146</u>	9	HPV	E7	8	A
QAEQATSNY	147	9	HPV	E7	46	
ATSNYYIVTY	<u>148</u>	10	HPV	E7	50	
ATDNYYIVTY	<u>149</u>	10	HPV	E7	50	A
RVLPPNWKY	<u>150</u>	9	Human	40s riboprot S13	132	
RLAHEVGWKY	<u>151</u>	10	Human	60s ribo prot L13A	139	
AYKKQFSQY	<u>152</u>	9	Human	60s ribo prot L5	217	
AADNPPAQY	<u>153</u>	9	Human	CEA	261	Α
RSGPGPGNVLY	154	11	Human	CEA	225	A
RSDGPGPGVLY	<u>155</u>	11	Human	CEA	225	Α
RSDSGPGPGLY	<u>156</u>	11	Human	CEA	225	Α
RSDSVGPGPGY	<u>157</u>	11	Human	CEA	225	Α
SLFVSNHAY	<u>158</u>	9	Human	fructose biphosphatealdolas e	355	
RWGLLLALL	159	9	Human	Her2/neu	8	
YTGPGPGVY	160	9	Human	Jchain	102	Α
YTAGPGPGY	161	9	Human	Jchain	102	A
TODLVOEKY	162	9	Human	MAGE1	240	
TOGPGPGKY	163	9	Human	MAGEI	240	Α
TQDGPGPGY	164	9	Human	MAGE1	240	A
EVGPGPGLY	165	9	Human	MAGE3	161	Α
EVDGPGPGY	166	9	Human	MAGE3	161	Α
IYGPGPGLIF	167	10	Human	MAGE3	195	A
RISGVDRYY	168	9	Human	NADH ubiqoxidoreductas	53	
IMVLSFLF	169	8	Pf	e CSP	427	
ALFQEYQCY	170	9	Pf	CSP	18	
LSEYYDXDIY	171	10	Pf		347	
FQAAESNERY	172	10	Pf		13	
ELEASISGKY	173	10	Pf		81	
FVSSIFISFY	174	10	Pf		255	
KVSDEIWNY	175	9	Pf		182	
IMNHLMTLY	175 176	9	Pf		38	
LIENELMNY	170 177	9	Pf		149	
NVDQQNDMY	178	9	Pf		182	
SSFFMNRFY	178	9	Pf		309	
OAAESNERY	180	9	Pf		14	
VAVESIVEY I	100	9	**		14	

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HLA-A1 SUPERTYPE

	SEQ ID				
Sequence	NO.	AA	Organism	Protein	Position Analog
LEASISGKY	181	9	Pf		82
NLALLYGEY	182	9	Pf		188
SSPLFNNFY	183	9	Pf		14
QNADKNFLY	<u>184</u>	9	Pf		145
VSSIFISFY	185	9	Pf		256
SYKSSKRDKF	<u>186</u>	10	Pf		225
RYQDPQNYEL	187	10	Pf		21
DFFLKSKFNI	188	10	Pf		3
NYMKIMNHL	189	9	Pf		34
TYKKKNNHI	190	9	Pf		264
SFFMNRFYI	191	9	Pf		310
FYITTRYKY	192	9	Pf		316
KYINFINFI	193	9	Pf		328
TWKPTIFLL	194	9	Pf		135
KYNYFIHFF	195	9	Pf		216
HFFTWGTMF	196	9	Pf		222
RMTSLKNEL	197	9	Pf		61
YYNNFNNNY	198	9	Pf		77
GTDEXRNXY	199	9	Unknown	Naturally	A
				processed	Α
ETDXXXDRSEY	200	11	Unknown	Naturally processed	A
FTDVNSXXRY	201	10	Unknown	Naturally	Α
MADDANAM	202	9	Unknown	processed Naturally	Α
VXDPYNXKY	202	9	Unknown	processed	A
VADKVHXMY	203	9	Unknown	Naturally	A
Description (20.4	9	Unknown	processed Naturally	Α
ETXXPDWSY	204	9	Onknown	processed	A
XTHNXVDXY	205	9	Unknown	Naturally	Α
				processed	

TABLE 12

HLA-A1 SUPERTYPE

SEQ ID							
Sequence	NO.	A*0101	A*2902	A*3002			
AYGPGPGKF	1		44854	3.2			
AEIPYLAKY	<u>2</u>			144			
AADAAAKY	<u>3</u>	20					
AYSSWMYSY	4			4.9			
LAEKTMKEY	<u>5</u>	174					
GTYDYWAGY	<u>6</u>	141					
LSVHSIONDY	2	279					
DTGQCPELVY	<u>8</u>	129					
DLLDTASALY	<u>9</u>		74	37			
WFHISCLTF	<u>10</u>	85324	95	75094			
LSLDVSAAFY	11	267	12	7.1			
LSGPGPGAFY	<u>12</u>	25	1383	6.6			
LSLGPGPGFY	<u>13</u>	21	132	8.2			
LSLDGPGPGY	14	266	274	181			
KTYGRKLHLY	<u>15</u>	171	27	1.5			
KTGPGPGHLY	16	29	192	1.3			
KTYGPGPGLY	<u>17</u>	5.7	227	0.96			
KTYGGPGPGY	18	282	228	1.7			
KYTSFPWL	<u>19</u>		>172413	346			
FAAPFTQCGY	20		461	1364			
SYOHFRKLLL	21	>83333	28	3768			
LYSHPIILGF	22	3166	109	1116			
MSTTDLEAY	23	0.00	2565	396			
MYVGGPGPGVF	24		89	2870			
VMGSSYGF	<u>25</u>		145	41967			
EVDGVRLHRY	<u>26</u>		14940	113			
RTEILDLWVY	<u>27</u>	99	10204	315			
RQDILDLWVY	<u>28</u>	8995	13928	95			
RTDILDLWVY	<u>29</u>	85	13424	360			
YTDGPGIRY	<u>30</u>	11	562	7911			
ATELHPEYY	<u>31</u>	43	6608	1734			
DLWVYHTQGYY	<u>32</u>	5880	852	16			
WVYHTQGYY	<u>33</u>	703	215	5.6			
FFLKEKGGF	<u>34</u>		3015	141			
LYVYHTQGY	<u>35</u>		216	258			
ITKILYQSNPY	<u>36</u>	>10060	64908	298			
KTLYQSNPY	<u>37</u>	6912	1703	35			
PVDPNLEPY	38	195	13193	7121			
STVKHHMY	<u>39</u>	8132	1760	68			
LSKISEYRHY	40	14306	55190	186			
ISEYRHYNY	<u>41</u>	25	1329	32			
RFHNIRGRW	<u>42</u>	52917	18	58			
RFLSKISEY	<u>43</u>	>40322	34623	23			
RFHNISGRW	44	48564	174	37			

HLA-A1 SUPERTYPE

	SEQ ID	CILICIA		
Sequence	NO.	A*0101	A*2902	A*3002
TLEKLTNTGLY	<u>45</u>	23	991	92
TLGPGPGTGLY	<u>46</u>	350	1320	7.4
TLEGPGPGGLY	<u>47</u>	11	2320	40
TLEKGPGPGLY	<u>48</u>	13	2036	40
TLEKLGPGPGY	49	269	4473	1962
TLEKLTNTGLY	<u>50</u>	77	5500	154
TLEKITNTELY	<u>51</u>	17	8402	3897
PYGVCIMCLRF	<u>52</u>		69	43722
ITDIILECVY	<u>53</u>	1.8	7660	505
YSDISEYRHY	<u>54</u>	3.8	1350	514
LTDIEITCVY	<u>55</u>	12	540	80
YSDIRELRHY	<u>56</u>	14	1137	740
ELSSALEIPY	<u>57</u>	171	6031	4472
ETSSALEIPY	<u>58</u>	19	12026	7144
ELDSALEIPY	<u>59</u>	38	82189	38284
YTKVSEFRWY	<u>60</u>	276	3308	420
YSDVSEFRWY	<u>61</u>	3.9	1842	1026
LTDVSIACVY	<u>62</u>	2.9	764	72
FTSRIRELRY	<u>63</u>	4.4	77	50
YSDIRELRYY	<u>64</u>	9.4	733	456
LTDLRLSCVY	<u>65</u>	45	1783	613
FTSKVRKYRY	<u>66</u>	64	6677	52
YSDVRKYRYY	<u>67</u>	19	849	794
FYSKVSEFRF	<u>68</u>		79	18453
FYSRIRELRF	<u>69</u>		83	12598
PYAVCRVCLF	<u>70</u>		407	5226
ITEYRHYNY	<u>71</u>	114	625	418
ISDYRHYNY	<u>72</u>	16	45	455
ITEYRHYQY	<u>73</u>	90	1030	526
ISDYRHYQY	<u>74</u>	13	37	382
LTDLLIRCY	<u>75</u>	13	6857	5515
KTDQRSEVY	<u>76</u>	84	200429	1174
AYRDLCIVY	77		7117	66
KYYSKISEY	<u>78</u>		702	1.3
KFYSKISEF	<u>79</u>		73339	306
RYHNIRGRW	80		122644	15
RFHNIRGRF	<u>81</u>		346	0.69
AYKDLFVVY	<u>82</u>		639	1.3
LFVVYRDSF	<u>83</u>		919	18
RYHNIAGHY	84		138	0.93
RFHNIAGHF	<u>85</u>		635	1.4
VYGTTLEKF	<u>86</u>		75267	220
AYADLTVVY	<u>87</u>		136	9.3
AFADLTVVF	88		779	137
RYLSKISEY	89		4247	1.1
RYHNISGRW	<u>90</u>		104884	13

HLA-A1 SUPERTYPE

	SEQ ID								
Sequence	NO.	A*0101	A*2902	A*3002					
AYKDLCIVY	91		5205	29					
RYHSIAGOY	92		544	1.4					
RFHSIAGQF	93		481	1.2					
KYLFTDLRI	94		78575	339					
KFLFTDLRF	9 <u>5</u>		44	152					
LYTDLRIVY	96		4.8	2.1					
LFTDLRIVF	97		164	2649					
RFLSKISEF	98		40103	2049					
	99		13707	430					
EYRHYQYSF	100			7.1					
RYHNIMGRW	101		106990	1.3					
RFHNIMGRF	101		174						
NFACTELKF			46	6826					
PYAVCRVCF	103		5602	316					
LYYSKVRKY	104		1452	28					
VYADLRIVY	105		8.2	8.3					
VFADLRIVF	<u>106</u>		87	24062					
NYSLYGDTF	<u>107</u>		20945	64					
RFHNISGRF	108		572	2.8					
FTDLTIVY	<u>109</u>	16	1275	39043					
FTDLRIVY	<u>110</u>	26	813	8060					
TLEKLTNTGLY	111	174							
LTDIEITCVY	112	33							
LTDVSIACVY	113	57							
ITDIILECVY	114	187							
KTDQRSEVY	115	41							
FTDLTIVY	116	34							
YSDIRELRYY	117	20							
YTKVSEFRWY	118	204							
FTSRIRELRY	119	25							
FTSKVRKYRY	120	37							
ISDYRHYNY	121	28							
ISEYRHYQY	122	40							
ISDYRHYOY	123	28							
EYRHYCYSLY	124	125	198	3.7					
EYRHYNYSLY	125	111027	956	12					
LTDLLIRCY	126	64	750						
ETRHYCYSLY	127	43	755	10					
EYDHYCYSLY	128	110081	799	77					
KTRYYDYSVY	129	2957	87841	0.71					
KYDYYDYSVY	130	186339	5749	11					
ETRHYNYSLY	131	445	5464	29					
EYDHYNYSLY	132	11251	777	93					
	133	195	805	408					
PTLKEYVLDLY	134	20	1509	408 54					
HTDTPTLHEY	135								
RTETPTLQDY	136	11	1987	239					
ETDPVDLLCY	130	6.4	4110	52640					

HLA-A1 SUPERTYPE

	OFO ID								
Sequence	SEQ ID NO.	A*0101	A*2902	A*3002					
QTEQATSNYY	137	11	9576	500					
ATDNYYIVTY	138	7.4	1918	65					
LTEYVLDLY	139	6.0	941	81					
QTEQATSNY	140	14	119081	3247					
ROAKOHTCY	141	>135135	155246	108					
RTAKOHTCY	142	5647	130343	346					
HTDTPTLHEY	143	3047	130343	340					
RTETPTLODY	144	40							
PTLKEYVLDLY	145	426							
LTEYVLDLY	146	8.0							
OAEOATSNY	147	132							
ATSNYYIVTY	148	428							
ATDNYYIVTY	149	19							
	150	19		3.0					
RVLPPNWKY	151			3.8					
RLAHEVGWKY AYKKOFSOY	152			5.8					
	153	0.2		3.3					
AADNPPAQY	154	9.2	11070						
RSGPGPGNVLY		172	11270	6.3					
RSDGPGPGVLY	<u>155</u> 156	12	13162	12					
RSDSGPGPGLY		3.3	11856	4.2					
RSDSVGPGPGY	<u>157</u>	23	31193	33					
SLFVSNHAY	158 150			1.1					
RWGLLLALL	<u>159</u>		61253	300					
YTGPGPGVY	160	2.7	2015	6.4					
YTAGPGPGY	<u>161</u>	7.0	28	755					
TQDLVQEKY	<u>162</u>	57	33304	3796					
TQGPGPGKY	163	4192	36746	3.2					
TQDGPGPGY	164	381	37093	541					
EVGPGPGLY	<u>165</u>	50	18183	45					
EVDGPGPGY	<u>166</u>	29	25775	5766					
IYGPGPGLIF	<u>167</u>		58	6845					
RISGVDRYY	<u>168</u>			3.0					
IMVLSFLF	169		111	30000					
ALFQEYQCY	<u>170</u>	>42016	149	1032					
LSEYYDXDIY	<u>171</u>	11	1647	489					
FQAAESNERY	<u>172</u>	8958	1780	372					
ELEASISGKY	<u>173</u>	142	21934	463					
FVSSIFISFY	<u>174</u>	118	22	84					
KVSDEIWNY	<u>175</u>	435	230	1.9					
IMNHLMTLY	<u>176</u>	150	1.7	1.8					
LIENELMNY	<u>177</u>	412	3936	169					
NVDQQNDMY	<u>178</u>	47	22173	79057					
SSFFMNRFY	<u>179</u>	239	36	7.5					
QAAESNERY	180	353	24281	3011					
LEASISGKY	<u>181</u>	57792	17824	87					
NLALLYGEY	182	275	138	102					

HLA-A1 SUPERTYPE

	SEQ ID			
Sequence	NO.	A*0101	A*2902	A*3002
SSPLFNNFY	183	117	389	73
QNADKNFLY	184	3811	24	663
VSSIFISFY	<u>185</u>	144	1800	55
SYKSSKRDKF	186		12594	88
RYQDPQNYEL	187		79717	189
DFFLKSKFNI	188		47714	491
NYMKIMNHL	189		45443	110
TYKKKNNHI	<u>190</u>		21642	162
SFFMNRFYI	<u>191</u>		200	1022
FYITTRYKY	<u>192</u>		9.6	7.5
KYINFINFI	<u>193</u>		25475	55
TWKPTIFLL	<u>194</u>		21155	306
KYNYFIHFF	<u>195</u>		319	2.7
HFFTWGTMF	<u>196</u>		4.0	220
RMTSLKNEL	<u>197</u>		40270	14
YYNNFNNNY	198		19	34
GTDEXRNXY	199	0.67		
ETDXXXDRSEY	<u>200</u>	2.0		
FTDVNSXXRY	201	0.20		
VXDPYNXKY	202	2.3		
VADKVHXMY	203	2.4		
ETXXPDWSY	204	11		
XTHNXVDXY	205	1.4		

TABLE 13

HLA-A2 SUPERTYPE

Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
FPFKYAAAV	206	9	Artificial sequence			Α
AMAKAAAAV	207	9	Artificial sequence			PolyA
AMAKAAAAL	208	9	Artificial sequence			PolyA
AMAKAAAAT	209	9	Artificial sequence			PolyA
AXAKAAAAL	210	9	Artificial sequence			PolyA
FVYGGSKTSL	211	10	EBNA		508	
ILGPGPGL	212	8	Flu	Ml	59	Α
GILGFVFTL	213	9	Flu	M1	58	
GLIYNRMGAV	214	10	Flu A	MI	129	
VLMEWLKTRPI	215	11	Flu A	MI	41	
FLPSDYFPSV	216	10	HBV	Core	18	Α
FLGPGPGPSV	217	10	HBV	core	18	Α
FLPGPGPGSV	218	10	HBV	core	18	Α
FLPSGPGPGV	219	10	HBV	core	18	Α
WLGPGPGFV	220	9	HBV	env	335	Α
WLSGPGPGV	221	9	HBV	env	335	Α
GVLGWSPQV	222	9	HBV	env	62	Α
PVLPIFFCV	223	9	HBV	env	377	Α
VVQAGFFLV	224	9	HBV	env	177	Α
FLLAQFTSA1	225	10	HBV	Pol	503	
YLLTLWKAGI	226	10	HBV	pol	147	
YLGPGPGAGI	227	10	HBV	pol	147	Α
YLLGPGPGGI	228	10	HBV	pol	147	Α
YLLTGPGPGI	229	10	HBV	pol	147	Α
HVYSHPIIV	230	9	HBV	pol	1076	Α
FVLSLGIHV	231	9	HBV	pol	562	Α
YVDDVVLGV	232	9	HBV	pol	538	Α
IVRGTSFVYV	233	10	HBV	pol	773	Α
SLGPGPGIAV	234	10	HIV	env	814	Α
SLLGPGPGAV	235	10	HIV	env	814	Α
SLLNGPGPGV	236	10	HIV	env	814	Α
KITPLCVTL	237	9	HIV	Env	134	Α
KLTPLCVTM	238	9	HIV	Env	134	Α
KLTPLCVPL	239	9	HIV	Env	134	Α
KLTPLCVSL	240	9	HIV	Env	134	Α
KLTPLCITL	241	9 .	HIV	Env	134	Α
QLTPLCVTL	242	9	HIV	Env	134	Α
KLTPRCVTL	242	9	HIV	Env	134	A
ELTPLCVTL	243	9	HIV	Env	134	Α
OMTFLCVOM	245	9	HIV	Env	134	Α
KMTFLCVQM	245	9	HIV	Env	134	A
KLTPLCVAL	246 247	9	HIV	Env	134	A
KLTPFCVTL	247 248	ý	HIV	Env	134	A
SLYNTVATL	248 249	9	HIV	GAG	77	-
VLAEAMSQT	249 250	9	HIV	Gag	386	Α
VLAEAMSQA		9	HIV	Gag	386	A
VLADAINIOUA	<u>251</u>	,	111 Y	Gag	200	^

HLA-A2 SUPERTYPE

Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
VLAEAMSQI	252	9	HIV	Gag	386	A
ILAEAMSQV	253	9	HIV	Gag	386	Α
VLAEAMSKV	254	9	HIV	Gag	386	A
VLAEAMSHA	255	9	HIV	Gag	386	A
ILAEAMSOA	256	9	HIV	Gag	386	Α
VLAEAMSRA	250 257	9	HIV	Gag	386	Α
VLAEAMATA	258	9	HIV	Gag	386	Α
ILAEAMASA	259	9	HIV	Gag	386	Α
MTHNPPIPV	260	9	HIV	Gag	271	Α
MTNNPPVPV	261	9	HIV	Gag	271	Α
MTSNPPIPV	262	9	HIV	Gag	271	A
MTSNPPVPV		9	HIV	Gag	271	A
MTSDPPIPV	<u>263</u>	9	HIV	Gag	271	A
MTGNPPIPV	264	9	HIV	Gag	271	A
MTGNPPVPV	<u>265</u>	9	HIV	Gag	271	A
	<u>266</u>	9	HIV	-	271	A
MTGNPAIPV	<u>267</u>	9	HIV	Gag	271	A
MTGNPSIPV	<u> 268</u>	9		Gag	271	A
MTANPPVPV	<u> 269</u>		ні∨	Gag		A
SLYNTVATL	<u>270</u>	9	hiv	gag	77	
QAHCNISRA	<u>271</u>	9	HIV	gp160	332	
FLKEKGGV	<u>272</u>	8	HIV	NEF	117	A
GLGAVSRDL	<u>273</u>	9	HIV	NEF	45	Α
GLITSSNTA	274	9	HIV	NEF	62	A
ALEEEEVGFPV	<u>275</u>	11	HIV	NEF	83	A
FLKEKGGLEGV	<u>276</u>	11	HIV	NEF	117	Α
FLKEKGGLDGV	277	11	HIV	NEF	117	Α
GLIYSKKRQEV	278	11	HIV	NEF	173	Α
LLYSKKRQEI	279	10	HIV	NEF	174	A
LLYSKKRQEIL	280	11	HIV	NEF	174	Α
RLDILDLWV	281	9	HIV	NEF	182	Α
EILDLWVYHV	282	10	HIV	NEF	185	Α
ILDLWVYHV	283	9	HIV	NEF	186	Α
ILDLWVYNV	284	9	HIV	NEF	186	Α
WLNYTPGPGT	285	10	HIV	NEF	204	Α
WQNYTPGPGV	286	10	HIV	NEF	204	Α
WLNYTPGPGI	287	10	HIV	NEF	204	Α
YLPGPGIRYPL	288	11	HIV	NEF	207	Α
YTPGPGIRYPV	289	11	HIV	NEF	207	Α
LLFGWCFKL	289 290	9	HIV	NEF	221	A
LTFGWCFKV	290 291	9	HIV	NEF	221	A
LLFGWCFKLV	291 292	10	HIV	NEF	221	A
FGVRPQVPL	_	9	HIV	nef	84	A
FTVRPQVPL	<u>293</u>	9	HIV	nef	84	A
FSVRPQVPL	<u>294</u>	9	HIV	nef	84	A
YLKEPVHGV	<u>295</u>	9	HIV	pol	476	A
	<u>296</u>	9	HIV	•	476 476	А
FLKEPVHGV	<u>297</u>			pol		
PVPLQLPPV	<u>298</u>	9	HIV	REV	74	A
LQLPPLERV	<u>299</u>	9	HIV	REV	77	A
LLLPPLERLTL	<u>300</u>	11	HIV	REV	77	A

HLA-A2 SUPERTYPE

Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
LQLPPLERLTV	301	11	HIV	REV	77	Α
ILWQVDRM	302	8	HIV	VIF	9	Α
KLGSLQYL	303	8	HIV	VIF	146	Α
KVGSLQYV	304	8	HIV	VIF	146	Α
TLHDLCQAV	305	9	HPV	E6	11	Α
TLQDIVLHL	306	9	HPV	E7	7	
TLGPGPGHL	307	9	HPV	E7	7	Α
TLQGPGPGL	308	9	HPV	E7	7	Α
TLSFVCPWCV	309	10	HPV	E7	94	Α
TLSFVCPWCA	310	10	HPV18	E7	93	
RTLHDLCQA	311	9	HPV33	E6	10	
TLHDLCQAL	312	9	HPV33	E6	11	
YLSGADLNL	313	9	Human	CEA	605	Α
YLEPGPVTA	314	9	Human	gp100	280	
LLDGTATLRL		10	Human	gp100	457	
KVYGLSAFV	315 316	9	Human	Her2/neu	369	Α
IISAVVAIL	316	9	Human	Her2/neu	654	A
ILSAVVGIL	317	9	Human	Her2/neu	654	A
IISAVVGFL	318	9	Human	Her2/neu	654	A
IISAVVGIV	319 320	9	Human	Her2/neu	654	A
KISAVVGIL	320	9	Human	Her2/neu	369	A
KIFAVVGIL	<u>321</u>	9	Human	Her2/neu	369	Ā
KIFASVAIL	322	9	Human	Her2/neu	369	A
ELVSEFSRV	323	9	Human	Her2/neu	971	Â
VLVHPOWVV	<u>324</u>	9	Human	Kallikrein2	53	A
VLVHPQWVLTV	325	11	Human	Kallikrein2	53	A
DLMLLRLSEPV	<u>326</u>	11	Human	Kallikrein2	120	A
PLVCNGVLQGV	<u>327</u>	11	Human	Kallikrein2	216	A
VLVHPQWVLTV	328	11	Human	Kallikrein2	53	A
PLVCNGVLQGV	<u>329</u>	11	Human	Kallikrein2	216	A
QLGPGPGLMEV	330	11	Human	MAGE3	159	Ā
QLVGPGPGMEV	<u>331</u>	11	Human	MAGE3	159	Ā
QLVFGPGPGEV	332	11	Human	MAGE3	159	Ā
QLVFGGPGPGV	333	11	Human	MAGE3	159	Â
ALGIGILTY	334	9	Human	MARTI	27	Ā
AMGIGILTV	<u>335</u>	9	Human	MARTI	27	Â
LLWQPIPV	<u>336</u>	8	Human	PAP	136	Λ.
LLGPGPGV	<u>337</u>	8	Human	PAP	136	Α
VLAKELKFVTL	338	11	Human	PAP	30	A
VLGPGPGFVTL	339	11	Human	PAP	30	Α
VLAGPGPGVTL	<u>340</u>	11	Human	PAP	30	Â
VLAGPGPGTL	341	11	Human	PAP	30	Ā
VLAKEGPGPGL	342	11	Human	PAP	30	Â
TLMSAMTNV	343	9	Human	PAP	112	A
ILYSAHDTTV	344	10	Human	PAP	384	A
IVYSAHDTTV	345	10	ruman Human	PAP	284	A
VTAKELKFV	346	9	Human	PAP	30	A
ITYSAHDTTV	<u>347</u>	10	Human	PAP	284	A
SLSLGFLFV	348	9	Human	PAP	204	^
SESECTED V	<u>349</u>	,	Tuman	i Ai		

HLA-A2 SUPERTYPE

	SEQ ID					
Sequence	NO.	AA	Organism	Protein	Position	Analog
SLSLGFLFLV	350	10	Human	PAP		
LLALFPPEGV	351	10	Human	PAP		
LVALFPPEGV	352	10	Human	PAP		
ALFPPEGVSV	353	10	Human	PAP		
GLHGQDLFGV	354	10	Human	PAP		
LLPPYASCHV	355	10	Human	PAP		
LLWQPIPVHV	356	10	Human	PAP		
MLLRLSEPV	357	9	Human	PSA	118	Α
ALGTTCYV	358	8	Human	PSA	143	Α
VLRLFVCFLI	359	10	Pf		2	
FLIFHFFLFL	360	10	Pf		9	
LIFHFFLFLL	361	10	Pf		10	
FLFLLYILFL	362	10	Pf		15	
RLPVICSFLV	363	10	Pf		32	
VICSFLVFLV	364	10	Pf		35	
FLVFLVFSNV	365	10	Pf		39	
MMIMIKFMGV	366	10	Pf		62	
FLLYILFLV	367	9	Pf		17	
VICSFLVFL	368	9	Pf		35	
ATYGIIVPV	369	9	Pf		159	
KIYKIIIWI	370	9	Pf		9	
YMIKKLLKI	371	9	Pf		23	
LMTLYQIQV	372	9	Pf		42	
FMGVIYIMI	373	9	Pf		68	
FMNRFYITT	374	9	Pf		312	
YQDPQNYEL	375	9	Pf		22	
KTWKPTIFL	376	9	Pf		134	
LLNESNIFL	377	9	Pf		142	
FIHFFTWGT	378	9	Pf		220	
VLFLQMMNV	379	9	Pf		180	
NQMIFVSSI	380	9	Pf		251	
MIFVSSIFI	381	9	Pf		253	
SIFISFYLI	382	9	Pf		258	
RLFEESLGI	383	9	Pf		293	
ALWGFFPVL		9	Unknown	A2		Α
	<u>384</u>			alloepitope	100	
SVYDFFVWL	<u>385</u>	9		TRP2	180	
FAPGFFPYL	<u>386</u>	9				
QLFEDKYAL	387	9				
MLLSVPLLL	388	9				

TABLE 14

		HLA-A	SUPERT	YPE		
	SEQ					
Sequence	ID NO.	A*0201	A*0202	A*0203	A*0206	A*6802
FPFKYAAAV	206	71 0201	0202	0200		92
AMAKAAAAV	207	181	196	6.7	1485	177
AMAKAAAAL	208	413	123	3.7	18500	320
AMAKAAAAT	209	15143	12413	84	37000	>26666.67
AXAKAAAAL	210	>50000	469	3300	37000	>11428.57
FVYGGSKTSL	211	296				
ILGPGPGL	212	672	45	530	1262	56099
GILGFVFTL	213	1.0	10	236	2.1	1395
GLIYNRMGAV	214	317				
VLMEWLKTRPI	215	464				
FLPSDYFPSV	216	8.5	3.3	3.2	2.2	276
FLGPGPGPSV	217	17	0.80	2.5	55	286
FLPGPGPGSV	218	98	18	4.0	665	332
FLPSGPGPGV	219	21	1.2	3.4	64	40
WLGPGPGFV	220	171	4.1	2.2	530	293
WLSGPGPGV	221	220	2.5	12	885	24
GVLGWSPOV	222	22	157	389	28	9428
PVLPIFFCV	223	8.7	3136	14286	22	1814
VVQAGFFLV	224	440	79	2503	81	617
FLLAQFTSAI	225	65	1.9	4.8	148	533
YLLTLWKAGI	226	20	19	20	40	1388
YLGPGPGAGI	227	161	1.0	4.2	548	315
YLLGPGPGGI	228	180	12	3.3	89	2064
YLLTGPGPGI	229	42	15	59	60	5678
HVYSHPIIV	230	150	1923	14	1199	123
FVLSLGIHV	231	45	399	2817	131	112
YVDDVVLGV	232	18	14	70	16	354
IVRGTSFVYV	232	50000	5301	69	5398	1217
SLGPGPGIAV	234	1131	5.3	11	917	281
SLLGPGPGAV	235	95	17	2.6	642	795
SLLNGPGPGV	236	65	3.8	14	63	45
KITPLCVTL	237	461	36	528	59	883
KLTPLCVTM	238	340	3.6	143	197	6288
KLTPLCVPL	239	15	0.25	297	135	67
KLTPLCVSL	240	67	2.4	240	16	5947
KLTPLCITL	241	1.7	0.27	23	1.7	9155
QLTPLCVTL	242	64	1.5	57	368	933
KLTPRCVTL	243	597	150	20	1554	>63492.06
ELTPLCVTL	244	7190	38	231	1919	32
QMTFLCVQM	245	3153	40	1127	232	1297
KMTFLCVQM	246	1793	22	525	100	8744
KLTPLCVAL	247	209	2.3	54	11	13009
KLTPFCVTL	248	87	0.37	28	78	11814
SLYNTVATL	249	290	6573	68	37000	20000
VLAEAMSOT	250	290	2.2	0.65	236	447
	250					

		HLA-A2	SUPERT	YPE		
	SEQ					
Sequence	ID NO.	A*0201	A*0202	A*0203	A*0206	A*6802
VLAEAMSOA	251	24	1.1	0.30	9.6	271
VLAEAMSOI	252	71	0.15	0.87	70	207
ILAEAMSQV	253	38	1.1	1.1	101	34
VLAEAMSKV	254	230	1.8	1.4	93	329
VLAEAMSHA	255	149	1.7	1.2	121	431
ILAEAMSQA	256	29	1.0	1.1	8.6	253
VLAEAMSRA	257	127	0.88	1.0	20	229
VLAEAMATA	258	6.7	1.4	0.73	8.6	33
ILAEAMASA	259	22	0.72	0.82	6.8	343
MTHNPPIPV	260	167	119	1.4	158	1.4
MTNNPPVPV	261	86	18	0.42	287	309
MTSNPPIPV	262	53	16	0.39	250	3.8
MTSNPPVPV	263	22	29	0.80	81	1.1
MTSDPPIPV	264	107	13	0.45	587	2.5
MTGNPPIPV	265	125	11	0.74	79	7.8
MTGNPPVPV	266	2021	158	23	35	0.84
MTGNPAIPV	267	1200	24	10	213	0.48
MTGNPSIPV	268	16	1.1	0.43	257	0.57
MTANPPVPV		20	5.0	0.62	134	4.0
SLYNTVATL	269 270	367	79	19	15072	247113
QAHCNISRA	270	338	"	17	13072	24/113
FLKEKGGV	271	13327	653	267	>14341.09	>19464.72
GLGAVSRDL		18679	436	1733	>10393.26	>16666.67
GLITSSNTA	273 274	5800	102	64	7865	>14311.27
ALEEEEVGFPV	274 275	2420	487	15744	2988	>13793.1
FLKEKGGLEGV		322	3.5	6.8	739	1252
FLKEKGGLDGV	<u>276</u> 277	332	3.7	11	3207	3807
GLIYSKKROEV		8971	57	152	>8564.81	>14260.25
LLYSKKRQEI	278	80687	382	152	>9438.78	>15686.27
LLYSKKRQEIL	<u>279</u>	>38167.9	282	1569	>8564.81	>14260.25
LLISKKKQEIL	280	4	202	1309	×6504.61	~14200.23
RLDILDLWV	281	43	615	1639	2635	>17777.78
EILDLWVYHV	282	496	569	1865	2229	163
ILDLWVYHV	283	17	30	156	145	7414
ILDLWVYNV	284	40	30	201	135	5814
WLNYTPGPGT	285	547	124	231	>31623.93	11808
WQNYTPGPGV	286	1175	114	230	223	11993
WLNYTPGPGI	287	135	4.6	46	>31623.93	1196
YLPGPGIRYPL	288	1026	20	1583	3497	782
YTPGPGIRYPV	289	7764	1985	11126	1112	9.2
LLFGWCFKL	290	18	4.1	198	340	1084
LTFGWCFKV	291	15	33	1168	187	9.7
LLFGWCFKLV	292	658	84	114	1669	3276
FGVRPQVPL	293					321
FTVRPQVPL	294					13
FSVRPQVPL	295					52
•	293 296	54	0.65	1.9	212	63
YLKEPVHGV						

	HLA-A2 SUPERTYPE									
	SEQ									
Sequence	ID NO.	A*0201	A*0202	A*0203	A*0206	A*6802				
PVPLQLPPV	298	10047	>7337.88	12595	81	>15625				
LOLPPLERV	299	7951	7705	13517	203	1786				
LLLPPLERLTL	300	34	2607	9010	45	>12779.55				
LOLPPLERLTV	301	159	4545	6270	52	>61068.7				
ILWQVDRM		1745	67	2998	11332	>19464.72				
KLGSLQYL	302	1862	14	298	9010	>19464.72				
KVGSLQYV	303 304	1650	441	703	1904	17480				
TLHDLCQAV	305	331	17	15	10585	2809				
TLODIVLHL		22	4.4	46	781	5088				
TLGPGPGHL	306	14974	35	66	12144	27910				
TLOGPGPGL	307	6248	62	951	9121	3809				
TLSFVCPWCV	308	786	123	370	4357	388				
TLSFVCPWCV	309	1611	221	521	27321	13228				
RTLHDLCQA	310	8121	34	678	96	61604				
	311	1404	2.7	40	2182	70390				
TLHDLCQAL YLSGADLNL	312	36	4.9	9.2	1605	51227				
YLEPGPVTA	<u>313</u>	36 466	10	27	20720	>470588.2				
YLEPGPVIA	314	400	10	21	20720	4/0300.2				
LLDGTATLRL	214	180	1.9	201	841	>421052.6				
	315					3				
KVYGLSAFV	316	33	1.8	11	69	110				
IISAVVAIL	317	1127	8.0	45	1440	148				
ILSAVVGIL	318	1464	1.9	21	2539	11854				
IISAVVGFL	319	747	1.0	4.8	234	77				
IISAVVGIV	320	712	15	20	958	390				
KISAVVGIL	321	6238	42	60	1752	4952				
KIFAVVGIL	322	3957	38	34	1539	6659				
KIFASVAIL	323	1062	16	21	1068	363				
ELVSEFSRV	324	8178	969	53	197	23				
VLVHPQWVV	325	464	65	1988	3224	14606				
VLVHPQWVLT	525	11	1.7	3.0	13	3288				
V	326									
DLMLLRLSEPV	327	69	66	32	118	2078				
PLVCNGVLQGV	328	91	424	36	212	3532				
VLVHPQWVLT		11	1.5	16	31	8889				
V	<u>329</u>									
PLVCNGVLQGV	<u>330</u>	26	126	19	264	4211				
QLGPGPGLMEV	<u>331</u>	194	9.4	29	481	648				
QLVGPGPGMEV	<u>332</u>	865	17	19	919	223				
QLVFGPGPGEV	<u>333</u>	2944	106	50	4067	447				
QLVFGGPGPGV	<u>334</u>	2153	96	242	3207	1318				
ALGIGILTV	<u>335</u>	11								
AMGIGILTV	<u>336</u>	15								
LLWQPIPV	337	137	2445	9.9	4251	32939				
LLGPGPGV	<u>338</u>	25	49	123	93	5620				
VLAKELKFVTL	339	1298	23	194	5170	15664				
VLGPGPGFVTL	340	1528	13	63	4766	42136				
VLAGPGPGVTL	341	1118	2.4	94	7200	2645				
VLAKGPGPGTL	<u>342</u>	11256	26	344	11450	>170212.7				

	HLA-A2 SUPERTYPE									
	SEQ									
Sequence	ID NO.	A*0201	A*0202	A*0203	A*0206	A*6802				
Sequence						7				
VLAKEGPGPGL	343	1890	6.9	37	59024	50993				
TLMSAMTNV	344	636	14	35	2188	484				
ILYSAHDTTV	345	397	1.1	13	1480	6285				
IVYSAHDTTV	346	7643	91	627	356	737				
VTAKELKFV	347	7143	2688	40	137	26667				
ITYSAHDTTV	348	4167	115	238	154	82				
SLSLGFLFV	349	77	25	21	93	26667				
SLSLGFLFLV	350	1.9	3.9	17	42	348				
LLALFPPEGV	351	5.0	0.73	1.6	148	163				
LVALFPPEGV	352	156	17	4.8	463	28				
ALFPPEGVSV	353	15	1.1	18	119	4444				
GLHGQDLFGV	354	12	2.3	3.1	18	>80000				
LLPPYASCHV	355	88	15	16	97	5333				
LLWQPIPVHV	356	25	1.8	18	285	62				
MLLRLSEPV	357	47	29	48	689	433				
ALGTTCYV	358	93	6.7	12	292	28284				
VLRLFVCFLI	_	2744	2112	299	68226	45639				
FLIFHFFLFL	359 360	161	174	2087	288	475				
LIFHFFLFLL		200	1468	3167	1562	460				
FLFLLYILFL	361	2834	172	2012	2113	8248				
RLPVICSFLV	362	12	2.5	33	19	9176				
VICSFLVFLV	363	167	415	2916	197	1949				
FLVFLVFSNV	<u>364</u>	269	212	35	232	5393				
MMIMIKFMGV	<u>365</u>	123	19	25	109	39				
FLLYILFLV	366	346	279	3091	1801	6981				
VICSFLVFL	<u>367</u>	184	19	2331	236	4800				
ATYGIIVPV	<u>368</u>	3.2	2.0	2.8	5.0	21				
KIYKIIIWI	<u>369</u>	157	1179	638	101	2198				
	<u>370</u>	105	4.6	4.7	93	63127				
YMIKKLLKI	<u>371</u>			20	615	1276				
LMTLYQIQV	<u>372</u>	14	1.6							
FMGVIYIMI	<u>373</u>	13	2.1	26	98 996	14501				
FMNRFYITT	<u>374</u>	101	18	13		6543				
YQDPQNYEL	<u>375</u>	79	18	441	52	166775				
KTWKPTIFL	<u>376</u>	135	1242	7487	76	3617				
LLNESNIFL	<u>377</u>	43	2.5	24	143	4484				
FIHFFTWGT	<u>378</u>	80	4.7	64	60	383				
VLFLQMMNV	<u>379</u>	31	1.8	2.7	9.5	323				
NQMIFVSSI	<u>380</u>	250	21	3.6	14	198				
MIFVSSIFI	381	85	18	83	114	5.2				
SIFISFYLI	382	289	35	1416	43	18				
RLFEESLGI	<u>383</u>	26	1.9	5.5	68	418				
ALWGFFPVL	<u>384</u>	3.6	0.74	3.7	15	1503				
SVYDFFVWL	385	36	169	226	10	0.86				
FAPGFFPYL	<u>386</u>	48	0.85	44	2.3	7.6				
QLFEDKYAL	<u>387</u>	646	1.8	380	2009	2982				
MLLSVPLLL	<u>388</u>	9.0	79	41	8.4	24607				

TABLE 15

HLA-A3 SUPERTYPE								
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog		
ALNAAAAAK		9	Artificial			Poly		
	389		sequence			•		
ALAAGAAAK		9	Artificial			Poly		
	<u>390</u>	_	sequence					
ALQAAAAK	201	9	Artificial			Poly		
STGPGPGVVRR	<u>391</u>	11	sequence HBV	core	141	Α		
STLGPGPGVRR	392	11	HBV	core	141	A		
STLPGPGPGRR	<u>393</u> 394	11	HBV	core	141	A		
STLPEGPGPGR		11	HBV	core	141	A		
OAGFFLLTR	<u>395</u>	9	HBV	ENV	179	^		
RVHFASPLH	<u>396</u>	9	HBV	POL	818			
AAYAAQGYK	397	9	HCV	II	1247			
KSKFGYGAK	398	9	HCV	II	2551			
PAAYAAQGYK	399	10	HCV	II	1246			
RMYVGGVEH	400	9	HCV	IV	635			
SOLSAPSLK	401	9	HCV	IV	2209			
TSCGNTLTCY	402	10	HCV	NS5	2740			
VTGPGPGPVWK	403	11	HIV	env	48	Α		
VTVGPGPGVWK	404	11	HIV	env	48	A		
VTVYGPGPGWK	405	11	HIV	env	48	A		
VTVYYGPGPGK	<u>406</u>	11	HIV	env	48	Ā		
	407	9	HIV	NEF	95	^		
PVRPQVPLR	408	10	HIV	NEF	61	Α		
HGAITSSNTK AVDLSFFLK	<u>409</u>	9	HIV	NEF	111	A		
	410	9			113	A		
DVSHFLKEK	411	-	HIV HIV	NEF NEF	113	A		
GVLDGLIYSK	412	10 9	HIV	NEF	124	A		
GVDGLIYSK	<u>413</u>	9			185	A		
EILDLWVYK	414	-	HIV	NEF				
ILDLWVYK	415	8	HIV	NEF	186	A		
RVPLTFGWCFK	<u>416</u>	11	HIV	NEF	216	A		
QVYTPGPGTR	<u>417</u>	10	HIV	NEF	205	A		
AVGPGPGLK	<u>418</u>	9	HIV	nef	84	A		
AVDGPGPGK	<u>419</u>	9	HIV	nef	84	A		
QMGPGPGNFK	<u>420</u>	10	HIV	pol	1432	A		
QMAGPGPGFK	<u>421</u>	10	HIV	pol	1432	A		
QMAVGPGPGK	<u>422</u>	10	HIV	pol	1432	A		
TVGPGPGPEK	<u>423</u>	10	HIV	pol	935	A		
TVQGPGPGEK	<u>424</u>	10	HIV	pol	935	A		
TVQPGPGPGK	<u>425</u>	10	HIV	pol	935	A		
VAIKIGGQLK	<u>426</u>	10	HIV	Pol	98	A		
VTVKIGGQLK	<u>427</u>	10	HIV	Pol	98	A		
VTIKVGGQLK	<u>428</u>	10	HIV	Pol	98	A		
VTIRIGGQLK	<u>429</u>	10	HIV	Pol	98	Α		
VTVRIGGQLK	<u>430</u>	10	HIV	Pol	98	Α		
VTVKVGGQLK	<u>431</u>	10	HIV	Pol	98	Α		
VTIRVGGQLK	<u>432</u>	10	HIV	Pol	98	Α		

HLA-A3 SUPERTYPE									
	SEQ ID								
Sequence	NO.	AA	Organism		Position				
VTVRVGGQLK	<u>433</u>	10	HIV	Pol		A .			
VTVKIGGQLR	<u>434</u>	10	HIV	Pol		A.			
VTIRIGGQLR	435	10	HIV	Pol		A			
VTIKLGGQIR	<u>436</u>	10	HIV	Pol		A.			
VSIKVGGQIK	437	10	HIV	Pol		A			
VSIRVGGQIK	438	10	HIV	Pol		A			
VTVKIEGQLK	439	10	HIV	Pol		A			
VTIKIEGQLK	440	10	HIV	Pol		A			
VTVKIEGQLR	441	10	HIV	Pol		A			
VSIRVGGQTK	442	10	HIV	Pol		A			
VSIRVGGQTR	443	10	HIV	Pol		A			
VTVRIGGMQK	444	10	HIV	Pol	98	A.			
ITVKIGKEVR	445	10	HIV	Pol	98 .	A.			
GTRQARRNK	446	9	HIV	REV	36	A.			
GTRQARRNRK	447	10	HIV	REV	36	A.			
GTRQARRNRRK	448	11	HIV	REV	36	A			
GTRQTRKNK	449	9	HIV	REV	37	A.			
GTRQTRKNRK	450	10	HIV	REV	37	A.			
GTRQTRKNRRK	451	11	HIV	REV	37	A			
RVRRRRWRAR	452	10	HIV	REV	43	A			
KVRRRRWRAR	453	10	HIV	REV	43	A.			
LTISYGRK	454	8	HIV	TAT	46	A.			
KTLGISYGR	455	9	HIV	TAT	44	A			
LTISYGRKK	456	9	HIV	TAT	46	A.			
GTSYGRKKR	457	9	HIV	TAT	47	A.			
GTGISYGRK	458	9	HIV	TAT	45	A.			
KTLGISYGRK	459	10	HIV	TAT	44	A			
LTISYGRKKR	460	10	HIV	TAT	46	A.			
KTLGISYGRKK	461	11	HIV	TAT	44	A			
TVCNNCYCK	462	9	HIV	TAT	23	Α			
LVISYGRKKRR	463	11	HIV	TAT	46	A			
ISYGRKKRROK	464	11	HIV	TAT	48	A.			
ETGPSGQPCK	465	10	HIV	TAT	101	A			
KVGPGGYPRR	466	10	HIV	TAT	101	A			
KAGPGGYPRK	467	10	HIV	TAT	101	A.			
KVGPGGYPRRK	468	11	HIV	TAT		A.			
AVPGGYPRR	469	9	HIV	TAT		A			
AVPGGYPRRK	470	10	HIV	TAT		A.			
KVGSLOYLK	471	9	HIV	VIF		A.			
ETVRHFPR	472	8	HIV	VPR		A			
AACHKCIDFY	473	10	HPV	E6	63	-			
LLIRCLRCQK	474	10	HPV	E6	101				
KISEYRHYNY	475	10	HPV	E6	72				
AVCRVCLLFY	476	10	HPV	E6	64				
FAFTDLTIVY	477 477	10	HPV	E6	45				
FAFADLTVVY	478	10	HPV	E6	45				
RFLSKISEYR	478 479	10	HPV	E6	68				
ILIRCIICOR	479 480	10	HPV	E6	99				
RTAMFQDPQER	481	11	HPV	E6	5				
	701				-				

HLA-A3 SUPERTYPE									
	SEQ ID								
Sequence	NO.	AA	Organism	Protein	Position Analo	g			
AMFQDPQERPR	482	11	HPV	E6	7				
MFQDPQERPRK	483	11	HPV	E6	8				
DLLIRCINCQK	484	11	HPV	E6	105				
RFEDPTRRPYK	485	11	HPV	E6	3				
ELTEVFEFAFK	486	11	HPV	E6	40				
GLYNLLIRCLR	487	11	HPV	E6	97				
NLLIRCLRCQK	488	11	HPV	E6	100				
EVLEESVHEIR	489	11	HPV	E6	17				
EVYKFLFTDLR	490	11	HPV	E6	41				
FLFTDLRIVYR	491	11	HPV	E6	45				
EVLEIPLIDLR	492	11	HPV	E6	20				
DLRLSCVYCKK	493	11	HPV	E6	28				
EVYNFACTELK	494	11	HPV	E6	44				
RVCLLFYSKVR	495	11	HPV	E6	67				
LLFYSKVRKYR	496	11	HPV	E6	70				
OLCDLLIRCYR	497	11	HPV	E6	98				
TLEQTVKK	498	8	HPV	E6	87				
ATRDLCIVYR	499	10	HPV	E6	53 A				
AFRDLCIVYK	500	10	HPV	E6	53 A				
ATCDKCLKFY	<u>500</u>	10	HPV	E6	68 A				
AVCDKCLKFR	502	10	HPV	E6	68 A				
KLYSKISEYR	503	10	HPV	E6	75 A				
KFYSKISEYK	<u>503</u>	10	HPV	E6	75 A				
KFSEYRHYCY	<u>504</u>	10	HPV	E6	79 A				
KISEYRHYCR	<u>505</u>	10	HPV	E6	79 A				
LFIRCINCQK	507	10	HPV	E6	106 A				
LLIRCINCOR	508	10	HPV	E6	106 A				
KVRFHNIRGR	<u>508</u>	10	HPV	E6	129 A				
KORFHNIRGK		10	HPV	E6	129 A				
WFGRCMSCCR	<u>510</u> 511	10	HPV	E6	139 A				
WTGRCMSCCK		10	HPV	E6	139 A				
MTCCRSSRTR	<u>512</u>	10	HPV	E6	144 A				
MSCCRSSRTK	<u>513</u>	10	HPV	E6	144 A				
STCRSSRTRR	<u>514</u>	10	HPV	E6	145 A				
SCCRSSRTRK	<u>515</u>	10	HPV	E6	145 A				
DIEITCVYCR	<u>516</u>	10	HPV	E6	27 A				
FTFKDLFVVY	<u>517</u>	10	HPV	E6	47 A				
FAFKDLFVVK	<u>518</u>	10	HPV	E6	47 A				
	<u>519</u>	10	HPV	E6	47 A 48 A				
AVKDLFVVYR	<u>520</u>								
AFKDLFVVYK	<u>521</u>	10	HPV	E6					
FVVYRDSIPK	<u>522</u>	10	HPV	E6	53 A				
DTIPHAACHK	<u>523</u>	10	HPV	E6	58 A				
DSIPHAACHR	<u>524</u>	10	HPV	E6	58 A				
KFIDFYSRIR	<u>525</u>	10	HPV	E6	67 A				
DTVYGDTLEK	<u>526</u>	10	HPV	E6	83 A				
DSVYGDTLER	<u>527</u>	10	HPV	E6	83 A				
LFIRCLRCQK	<u>528</u>	10	HPV	E6	101 A				
LLIRCLRCQR	<u>529</u>	10	HPV	E6	101 A				
RVHNIAGHYR	<u>530</u>	10	HPV	E6	126 A				

HLA-A3 SUPERTYPE								
	SEQ ID							
Sequence	NO.	AA	Organism	Protein	Position			
RFHNIAGHYK	<u>531</u>	10	HPV	E6	126	A		
RTQCHSCCNR	<u>532</u>	10	HPV	E6	135	Α		
RGQCHSCCNK	<u>533</u>	10	HPV	E6	135	Α		
ATTDLTIVYR	<u>534</u>	10	HPV	E6	46	Α		
AFTDLTIVYK	<u>535</u>	10	HPV	E6	46	Α		
RLYSKVSEFR	<u>536</u>	10	HPV	E6	68	Α		
RFYSKVSEFK	<u>537</u>	10	HPV	E6	68	Α		
KFSEFRWYRY	538	10	HPV	E6	72	Α		
KVSEFRWYRR	539	10	HPV	E6	72	Α		
YFVYGTTLEK	540	10	HPV	E6	81	Α		
YSVYGTTLER	541	10	HPV	E6	81	Α		
GTTLEKLTNR	542	10	HPV	E6	85	Α		
LVIRCITCQR	543	10	HPV	E6	99	Α		
LLIRCITCQK	544	10	HPV	E6	99	Α		
WVGRCIACWR	545	10	HPV	E6	132	Α		
WTGRCIACWK	546	10	HPV	E6	132	Α		
RTIACWRRPR	547	10	HPV	E6	135	Α		
RCIACWRRPK	548	10	HPV	E6	135	Α		
AVADLTVVYR	549	10	HPV	E6	46	Α		
AFADLTVVYK	550	10	HPV	E6	46	Α		
RVLSKISEYR	551	10	HPV	E6	68	Α		
RFLSKISEYK	552	10	HPV	E6	68	Α		
KFSEYRHYNY	553	10	HPV	E6	72	Α		
KISEYRHYNR	554	10	HPV	E6	72	Α		
ITIRCIICQR	555	10	HPV	E6	99	Α		
ILIRCIICOK	556	10	HPV	E6	99	Α		
WVGRCAACWR	557	10	HPV	E6	132	Α		
WAGRCAACWK	558	10	HPV	E6	132	Α		
CFACWRSRRR	559	10	HPV	E6	136	Α		
DTSIACVYCK	560	10	HPV	E6	27	Α		
DVSIACVYCR	<u>561</u>	10	HPV	E6	27	Α		
CVYCKATLEK	562	10	HPV	E6	32	A		
RFEVYQFAFK	563	10	HPV	E6	41	A		
RTEVYQFAFR	<u>564</u>	10	HPV	E6	41	A		
AVKDLCIVYR	565	10	HPV	E6	48	A		
AFKDLCIVYK	566	10	HPV	E6	48	A		
ATCHKCIDFY	<u>567</u>	10	HPV	E6	63	A		
AACHKCIDFK	<u> 568</u>	10	HPV	E6	63	A		
NLVYGETLEK	569	10	HPV	E6	83	A		
NSVYGETLER	<u> 570</u>	10	HPV	E6	83	A		
LSIRCLRCQK	570 571	10	HPV	E6	101	A		
LLIRCLRCQY		10	HPV	E6	101	A		
RVHSIAGQYR	<u>572</u>	10	HPV	E6	126	A		
RFHSIAGQYK	<u>573</u>	10	HPV	E6	126	A		
LVTDLRIVYR	<u>574</u>	10	HPV	E6	46	A		
	<u>575</u>	10	HPV	E6	46	A		
LFTDLRIVYK	<u>576</u>	10	HPV	E6	63	A		
CTMCLRFLSK	<u>577</u>	10	HPV	E6	63	A		
CIMCLRFLSR	<u>578</u>	10	HPV	E6	68	A		
RLLSKISEYR	<u>579</u>	10	nr v	EO	00	Α		

	HI	A-A3	SUPERTY	PE		
	SEQ ID					
Sequence	NO.	AA	Organism		Position	
RFLSKISEYY	<u>580</u>	10	HPV	E6	68	A
SFYGKTLEER	<u>581</u>	10	HPV	E6	82	Α
SLYGKTLEEK	<u>582</u>	10	HPV	E6	82	A
WFGRCSECWR	<u>583</u>	10	HPV	E6	132	Α
WTGRCSECWK	<u>584</u>	10	HPV	E6	132	Α
AFCRVCLLFY	<u>585</u>	10	HPV	E6	64	A
AVCRVCLLFR	<u>586</u>	10	HPV	E6	64	Α
CFLFYSKVRK	587	10	HPV	E6	69	Α
CLLFYSKVRR	<u>588</u>	10	HPV	E6	69	Α
LVYSKVRKYR	589	10	HPV	E6	71	Α
LFYSKVRKYK	590	10	HPV	E6	71	A
GTTLESITKK	591	10	HPV	E6	88	Α
WVGSCLGCWR	592	10	HPV	E6	135	Α
WTGSCLGCWK	593	10	HPV	E6	135	Α
VVADLRIVYR	594	10	HPV	E6	46	A
VFADLRIVYK	595	10	HPV	E6	46	Α
RTLSKISEYR	596	10	HPV	E6	68	Α
RLLSKISEYK	597	10	HPV	E6	68	Α
KVSEYRHYNY	598	10	HPV	E6	72	Α
KISEYRHYNK	599	10	HPV	E6	72	Α
IVIRCIICQR	600	10	HPV	E6	99	Α
WLGRCAVCWR	601	10	HPV	E6	132	A
WTGRCAVCWK	602	10	HPV	E6	132	Α
YVVCDKCLK	603	9	HPV	E6	67	A
YAVCDKCLR	604	9	HPV	E6	67	A
SVCRSSRTR	605	9	HPV	E6	145	A
SCCRSSRTK	606	9	HPV	E6	145	A
SLPHAACHK		9	HPV	E6	59	A
SIPHAACHR	607	9	HPV	E6	59	A
FVDLTIVYR	608	9	HPV	E6	47	A
FTDLTIVYK	609	9	HPV	E6	47	A
SFYGTTLEK	610	9	HPV	E6	82	A
SVYGTTLER	611	9	HPV	E6	82	A
TFLEKLTNK	612	9	HPV	E6	86	A
TTLEKLTNR	613	9	HPV	E6	86	A
ETNPFGICK	614	9	HPV	E6	56	A
	615	9	HPV	E6	56	A
EGNPFGICR	<u>616</u>	9	HPV	E6	86	A
NTLEQTVKR	<u>617</u>					A
ALCWRSRRR	<u>618</u>	9	HPV	E6	137	
AACWRSRRK	<u>619</u>	9	HPV	E6	137	A
VSIACVYCR	<u>620</u>	9	HPV	E6	28	A
SIACVYCKK	<u>621</u>	9	HPV	E6	29	A
ILYRDCIAY	<u>622</u>	9	HPV	E6	54	A
IVYRDCIAR	<u>623</u>	9	HPV	E6	54	A
CTAYAACHK	<u>624</u>	9	HPV	E6	59	A
CIAYAACHR	<u>625</u>	9	HPV	E6	59	A
SFYGETLEK	<u>626</u>	9	HPV	E6	84	A
SVYGETLER	<u>627</u>	9	HPV	E6	84	A
LIRCLRCQR	<u>628</u>	9	HPV	E6	102	Α

	H	LA-A3	SUPERTY	PE	
	SEQ ID				
Sequence	NO.	AA	Organism	Protein	Position Analog
RTQCVQCKK	<u>629</u>	9	HPV	E6	27 A
RLQCVQCKR	<u>630</u>	9	HPV	E6	27 A
KFLEERVKK	<u>631</u>	9	HPV	E6	86 A
KTLEERVKR	<u>632</u>	9	HPV	E6	86 A
NVMGRWTGR	633	9	HPV	E6	127 A
NIMGRWTGK	634	9	HPV	E6	127 A
LTYRDDFPY	635	9	HPV	E6	55 A
LVYRDDFPK	636	9	HPV	E6	55 A
RFCLLFYSK	637	9	HPV	E6	67 A
RVCLLFYSR	638	9	HPV	E6	67 A
LTFYSKVRK	639	9	HPV	E6	70 A
LLFYSKVRR	640	9	HPV	E6	70 A
ATLESITKR	641	9	HPV	E6	89 A
KVLCDLLIR	642	9	HPV	E6	97 A
KOLCDLLIK	643	9	HPV	E6	97 A
TFVHEIELK	644	9	HPV	E6	21 A
TSVHEIELR	645	9	HPV	E6	21 A
YTFVFADLR	646	9	HPV	E6	43 A
DFLEQTLKK	647	9	HPV	E6	86 A
DTLEOTLKR	648	9	HPV	E6	86 A
LVRCIICQR	649	ģ	HPV	E6	100 A
LIRCIICQK		9	HPV	E6	100 A
RVAVCWRPR	<u>650</u>	9	HPV	E6	135 A
RCAVCWRPK	651 652	9	HPV	E6	135 A
AFCWRPRRR	653	ģ	HPV	E6	137 A
AVCWRPRRK	_	9	HPV	E6	137 A
LSFVCPWCA	<u>654</u>	9	HPV	E7	94
TFCCKCDSTLR	655	11	HPV	E7	56
LVVESSADDLR	<u>656</u>	11	HPV	E7	74
	<u>657</u>	11	HPV	E7	88
TLQVVCPGCAR	658	11	HPV	E7	59
YLIHVPCCECK	<u>659</u>	11		E7	70
FVVQLDIQSTK	<u>660</u>		HPV		59
HTCNTTVR	<u>661</u>	8	HPV	E7	
GLVCPICSQK	<u>662</u>	10	HPV	E7	88 A
GFNHQHLPAR	<u>663</u>	10	HPV	E7	43 A
GVNHQHLPAK	<u>664</u>	10	HPV	E7	43 A
NVVTFCCQCK	<u>665</u>	10	HPV	E7	53 A
NIVTFCCQCR	<u>666</u>	10	HPV	E7	53 A
GVSHAQLPAK	<u>667</u>	10	HPV	E7	44 A
LIHVPCCECR	<u>668</u>	10	HPV	E7	60 A
AVLQDIVLH	<u>669</u>	9	HPV	E7	6 A
ATLQDIVLK	<u>670</u>	9	HPV	E7	6 A
GVNHQHLPK	<u>671</u>	9	HPV	E7	43 A
HVMLCMCCK	<u>672</u>	9	HPV	E7	59 A
HTMLCMCCR	<u>673</u>	9	HPV	E7	59 A
LSFVCPWCR	<u>674</u>	9	HPV	E7	94 A
AQPATADYK	675	9	HPV	E7	45 A
VVHAQLPAR	676	9	HPV	E7	45 A
VSHAQLPAK	677	9	HPV	E7	45 A

	н	A-A3	SUPERTY	PE	
	SEQ ID				
Sequence	NO.	AA	Organism		Position Analog
QLARQAKQH	678	9	HPV	E7	48 A
KQHTCYLIR	679	9	HPV	E7	54 A
VTLDIQSTK	<u>680</u>	9	HPV	E7	72 A
VQLDIQSTR	681	9	HPV	E7	72 A
SLGPGPGTK	682	9	Human	MAGE1	96 A
SLFGPGPGK	683	9	Human	MAGE1	96 A
LVGPGPGK	684	8	Human	MAGE2	116 A
KMFLQLAK	685	8	Human	p53	132
KMGPGPGK	686	8	Human	p53	132 A
KQENWYSLKK	687	10	Pf	CSP	58
GVGPGPGLK	688	9	Pf	LSA1	105 A
GVSGPGPGK	689	9	Pf	LSA1	105 A
FLLYILFLVK	690	10	Pf		17
LVFSNVLCFR	691	10	Pf		43
SSFDIKSEVK	692	10	Pf		116
TLYQIQVMKR	693	10	Pf		44
KQVQMMIMIK	694	10	Pf		58
GVIYIMIISK	695	10	Pf		70
ELFDKDTFFK	696	10	Pf		158
ALERLLSLKK	697	10	Pf		50
KILIKIPVTK		10	Pf		109
RLPLLPKTWK	698	10	Pf		128
SOVSNSDSYK	<u>699</u> 700	10	Pf		161
OONOESKIMK	_	10	Pf		197
IIALLIIPPK	<u>701</u>	10	Pf		249
SSPLFNNFYK	<u>702</u>	10	Pf		14
FLYLLNKKNK	<u>703</u>	10	Pf		151
LQMMNVNLQK	<u>704</u>	10	Pf		183
LTNHLINTPK	<u>705</u>	10	Pf		195
IFISFYLINK	<u>706</u>	10	Pf		259
RLFEESLGIR	<u>707</u>	10	Pf		293
LLYILFLVK	<u>708</u>	9	Pf		18
KSMLKELIK	709	9	Pf		129
	<u>710</u>				
PVLTSLFNK	<u>711</u>	9	Pf		166
KTMNNYMIK	<u>712</u>	9	Pf		18
LFDKDTFFK	<u>713</u>	9	Pf		159
YLFNQHIKK	<u>714</u>	9	Pf		287
MQSSFFMNR	<u>715</u>	9	Pf		307
RFYITTRYK	<u>716</u>	9	Pf		315
TTRYKYLNK	717	9	Pf		319
AVIFTPIYY	<u>718</u>	9	Pf		34
ALERLLSLK	<u>719</u>	9	Pf		50
SISGKYDIK	<u>720</u>	9	Pf		85
EQRLPLLPK	<u>721</u>	9	Pf		126
IALLIIPPK	722	9	Pf		250
PVVCSMEYK	<u>723</u>	9	Pf		270
VVCSMEYKK	<u>724</u>	9	Pf		271
FSYDLRLNK	<u>725</u>	9	Pf		308
HLNIPIGFK	<u>726</u>	9	Pf		323

	HLA-A3 SUPERTYPE						
Sequence	SEQ ID NO.	AA	Organism	Protein	Position Analog		
PLFNNFYKR	727	9	Pf		16		
YQNFQNADK	728	9	Pf		141		
QMMNVNLQK	729	9	Pf		184		
AVSEIQNNK	730	9	Pf		222		
GTMYILLKK	731	9	Pf		236		
FISFYLINK	732	9	Pf		260		
YLINKHWQR	733	9	Pf		264		
ALKISQLQK	734	9	Pf		273		
KINSNFLLK	735	9	Pf		282		
AAMXDPTTFK		10	Unknown	Naturally	Α		
	<u>736</u>			processed			
GTMTTSXYK		9	Unknown	Naturally	Α		
	737	•		processed			
SXXPAXFQK	720	9	Unknown	Naturally processed	A		
ATAGDGXXEXRK	<u>738</u>	12	Unknown	Naturally	Α		
ATAGDOXAEARK	<u>739</u>	12	CHAHOWH	processed	Α		

TABLE 16

		HLA-A3	SUPERTYPE			
	SEQ ID					
Sequence	NO.	A*0301	A*1101	A*3101	A*3301	A*6801
ALNAAAAK	389	74	21	10954	>72500	80000
ALAAGAAAK	390	19	37			
ALQAAAAK	<u>391</u>	57	65	51962	>72500	>80000
STGPGPGVVRR	<u>392</u>	18695	367	95	5983	5.8
STLGPGPGVRR	<u>393</u>	892	19	42	670	3.8
STLPGPGPGRR	<u>394</u>	297	19	61	1893	25
STLPEGPGPGR	<u>395</u>	325	26	28	822	30
QAGFFLLTR	<u>396</u>	10138	1678	302	182	5.3
RVHFASPLH	<u>397</u>	12	60	572	>122881.36	7620
AAYAAQGYK	398	18	18	1175	14074	34
KSKFGYGAK	<u>399</u>	36	596	116	>122881.36	>7626.31
PAAYAAQGYK	400	950	456	20314	>110687.02	666
RMYVGGVEH	401	3.8	274	162	>122881.36	>28776.98
SQLSAPSLK	402	306	25	1276	>122881.36	3845
TSCGNTLTCY	403	>36666.67	5.0			
VTGPGPGPVWK	404	2900	24	12964	>102836.88	425
VTVGPGPGVWK	405	174	2.7	2731	75360	21
VTVYGPGPGWK	406	1151	18	>8995.5	>102836.88	206
VTVYYGPGPGK	407	310	24	9720	101830	30
PVRPQVPLR	408	>10901.88	16112	332	3439	7012
HGAITSSNTK	409	2837	344	>16143.5	>22924.9	1235
AVDLSFFLK	410	226	23	6207	>27831.09	4038
DVSHFLKEK	411	>9298.39	5645	>17839.44	232	135
GVLDGLIYSK	412	1080	21	6007	>25151.78	831
GVDGLIYSK	413	10089	47	>17664.38	>29652.35	5100
EILDLWVYK	414	1032	64	>5774.78	288	93
ILDLWVYK	415	1265	320	13680	30096	12092
RVPLTFGWCFK	416	69	30	102	26651	571
QVYTPGPGTR	417	1249	852	1764	3334	273
AVGPGPGLK	418	18	3.6	128	75754	444
AVDGPGPGK	419	179	19	36837	>112403.1	2132
OMGPGPGNFK	420	49	22	2682	100771	63
QMAGPGPGFK	421	9.4	6.2	667	4784	30
QMAVGPGPGK	422	33	16	5961	86676	22
TVGPGPGPEK	423	115	17	10140	98177	23
TVQGPGPGEK	424	218	3.4	9874	103379	195
TVQPGPGPGK	425	41	2.5	1335	68584	28
VAIKIGGQLK	426	2593	151	46875	51222	123
VTVKIGGQLK	427	296	61	24385	104757	147
VTIKVGGQLK	428	188	59	6061	47647	127
VTIRIGGQLK	429	51	14	4458	65764	25
VTVRIGGQLK	430	226	15	5380	40344	49
VTVKVGGQLK	431	206	54	21484	46182	104
VTIRVGGOLK	432	43	13	3591	86086	28
VTVRVGGQLK	433	216	19	8238	>72319.2	141
VTVKIGGQLR	434	19185	194	417	3833	52

HLA-A3 SUPERTYPE							
Samue	SEQ ID NO.	A*0301	A*1101	A*3101	A*3301	A*6801	
Sequence VTIRIGGQLR	435	3192	23	61	1352	16	
VTIKLGGQLK VTIKLGGQIR		43252	219	590	12965	104	
VSIKVGGQIK	436	1921	86	57069	>72319.2	2026	
VSIR VGGQIK VSIR VGGQIK	437	642	91	50677	>61702.13	1960	
	438	647	23	4616	64604	30	
VTVKIEGQLK	<u>439</u>	361	69	5077	58024	27	
VTIKIEGQLK	<u>440</u>		143	394	4057	146	
VTVKIEGQLR	441	35612				290	
VSIRVGGQTK	442	341	21	29949	38958		
VSIRVGGQTR	<u>443</u>	18531	241	466	8595	288	
VTVRIGGMQK	<u>444</u>	54	13	2583	44425	155	
ITVKIGKEVR	445	>69182.39	12904	5057	24985	154	
GTRQARRNK	<u>446</u>	67	749	9713	45966	59708	
GTRQARRNRK	447	100	634	3800	>42335.77	7788	
GTRQARRNRRK	<u>448</u>	404	2596	7774	>24308.47	9104	
GTRQTRKNK	449	198	3104	13373	>29713.11	18657	
GTRQTRKNRK	<u>450</u>	129	1082	2485	60183	5998	
GTRQTRKNRRK	<u>451</u>	478	4184	4008	>24308.47	>17167.3	
RVRRRRWRAR	<u>452</u>	2443	>16759.78	265	3758	>36866.3	
KVRRRRWRAR	<u>453</u>	327	>20905.92	342	3243	15501	
LTISYGRK	454	988	708	27068	38162	482	
KTLGISYGR	455	53	9.8	21	502	36	
LTISYGRKK	456	584	69	13918	59654	63	
GTSYGRKKR	457	9965	5916	225	21588	5778	
GTGISYGRK	458	480	77	58102	>43740.57	7407	
KTLGISYGRK	459	36	79	841	42378	1629	
LTISYGRKKR	460	7161	1229	71	2515	33	
KTLGISYGRKK	461	52	285	91	23401	647	
TVCNNCYCK	462	9920	267	8793	28481	876	
LVISYGRKKRR	463	>11702.13	8669	562	267	4662	
ISYGRKKRRQK	464	48	2807	3147	>20000	4428	
ETGPSGOPCK	465	>14569.54	3501	>22500	>17813.27	50	
KVGPGGYPRR	466	2268	487	250	7904	721	
KAGPGGYPRK	467	62	43	10734	>17813.27	5555	
KVGPGGYPRRK	468	70	87	775	>5063.73	921	
AVPGGYPRR	469	3012	1215	1349	3453	109	
AVPGGYPRRK	470	819	60	39974	>5570.5	846	
KVGSLQYLK	471	482	70	2104	>43740.57	4200	
ETVRHFPR	472	>13513.51	4183	1000	81	86	
AACHKCIDFY	473	18824	261	20643	>116465.86	32548	
LLIRCLRCOK	474	437	170	6612	28936	78	
KISEYRHYNY	475	42	112	1426	35341	25077	
AVCRVCLLFY	475 476	77	21	1978	4520	1302	
FAFTDLTIVY		40343	21161	42065	131202	346	
FAFADLTVVY	477	18592	5866	23676	26768	402	
RFLSKISEYR	<u>478</u>	1640	18468	33	436	172	
KFLSKISE Y K ILIRCIICOR	479	8550	5012	33 377	2480	537	
ILIRCIICQR RTAMFQDPQER	<u>480</u>	8550 1478	103	49	2480 3459	19	
K LAWITODPOEK	481	14/8	103	49		19	

		HLA-A3	SUPERTYPE			
	SEQ ID					
Sequence	NO.	A*0301	A*1101	A*3101	A*3301	A*6801
MFQDPQERPRK	483	15493	8571	604	419	16729
DLLIRCINCQK	484	2923	935	4884	29	263
RFEDPTRRPYK	485	169	432	53	1758	7338
ELTEVFEFAFK	486	8966	582	25205	1733	15
GLYNLLIRCLR	487	1268	1568	250	401	1624
NLLIRCLRCQK	488	1565	854	3140	397	1480
EVLEESVHEIR	489	>45643.15	>20202.02	31037	212	240
EVYKFLFTDLR	490	31240	602	759	4.3	11
FLFTDLRIVYR	491	672	227	58	21	1.4
EVLEIPLIDLR	492	>47008.55	16638	36427	72	27
DLRLSCVYCKK	493	3644	1907	17023	109	3002
EVYNFACTELK	494	1622	117	484	5.9	2.7
RVCLLFYSKVR	495	771	190	221	1061	1267
LLFYSKVRKYR	496	28	94	7.0	11	15
QLCDLLIRCYR	497	1240	700	450	106	489
TLEQTVKK	498	4766	203	>100000	>75324.68	21400
ATRDLCIVYR	499	237	156	4.7	44	28
AFRDLCIVYK	500	31	15	10	132	57
ATCDKCLKFY	501	194	17	491	18080	4562
AVCDKCLKFR	502	77	15	11	45	34
KLYSKISEYR	503	5.4	168	6.4	28	91
KFYSKISEYK	504	7.6	674	27	329	208
KFSEYRHYCY	505	5092	7485	308	49397	14571
KISEYRHYCR	<u>506</u>	486	688	25	833	1488
LFIRCINCQK	<u>507</u>	2880	702	52	42	56
LLIRCINCQR	<u>508</u>	2818	686	30	50	14
KVRFHNIRGR	<u>509</u>	39	8632	27	4500	3979
KQRFHNIRGK	<u>510</u>	55	1953	573	35208	22879
WFGRCMSCCR	<u>511</u>	16071	10690	288	98	303
WTGRCMSCCK	<u>512</u>	6687	841	6496	15191	118
MTCCRSSRTR	<u>513</u>	3825	933	410	601	2.2
MSCCRSSRTK	<u>514</u>	352	169	2333	6916	12
STCRSSRTRR	<u>515</u>	2989	118	152	1020	312
SCCRSSRTRK	<u>516</u>	326	3272	5592	20916	8777
DIEITCVYCR	<u>517</u>	2014	826	3780	448	422
FTFKDLFVVY	<u>518</u>	14364	1208	10757	2725	62
FAFKDLFVVK	<u>519</u>	783	71	525	1066	3.6
AVKDLFVVYR	<u>520</u>	1728	91	3.1	9.1	3.3
AFKDLFVVYK	<u>521</u>	3256	211	32	93	576
FVVYRDSIPK	<u>522</u>	265	81	6216	146	30
DTIPHAACHK	<u>523</u>	2366	701	1763	9.3	23
DSIPHAACHR	<u>524</u>	2772	853	357	2.2	27
KFIDFYSRIR	<u>525</u>	8891	9008	3.3	677	2551
DTVYGDTLEK	<u>526</u>	50	15	28754	55090	31
DSVYGDTLER	<u>527</u>	292	23	485	891	28
LFIRCLRCQK	<u>528</u>	3390	1533	218	77	200
LLIRCLRCQR	<u>529</u>	3360	1396	28	75	13
RVHNIAGHYR	530	30	21	22	114	18

HLA-A3 SUPERTYPE							
Sequence	SEQ ID NO.	A*0301	A*1101	A*3101	A*3301	A*6801	
RFHNIAGHYK	531	25	22	2.6	80	23	
RTOCHSCCNR	531 532	338	20	22	132	161	
RGQCHSCCNK	532 533	6135	113	425	37669	20340	
ATTDLTIVYR	533 534	247	10	34	1739	14	
AFTDLTIVYK	534 535	701	112	3952	9380	215	
RLYSKVSEFR		6.4	131	24	690	73	
RFYSKVSEFK	<u>536</u>	27	521	30	4452	547	
KFSEFRWYRY	<u>537</u>	4750	1595	34	856	12811	
KVSEFRWYRR	<u>538</u>	266	16	2.8	159	30	
YFVYGTTLEK	<u>539</u>	204	62	2167	15740	53	
YSVYGTTLER YSVYGTTLER	<u>540</u>	430	96	2136	6903	19	
	<u>541</u>		1720	382	706	2946	
GTTLEKLTNR	<u>542</u>	3604		54	135	14	
LVIRCITCQR	<u>543</u>	2222	255	3009	2165	40	
LLIRCITCQK	<u>544</u>	291	120				
WVGRCIACWR	545	6227	1391	85	13	9.7	
WTGRCIACWK	<u>546</u>	2633	55	3078	169	24	
RTIACWRRPR	<u>547</u>	40	63	3.2	95	51	
RCIACWRRPK	<u>548</u>	1535	1476	292	176	1655	
AVADLTVVYR	<u>549</u>	489	11	31	892	7.3	
AFADLTVVYK	<u>550</u>	2365	107	1113	13557	50	
RVLSKISEYR	<u>551</u>	34	84	24	197	136	
RFLSKISEYK	<u>552</u>	31	287	42	10237	112	
KFSEYRHYNY	<u>553</u>	5819	5521	286	18351	1798	
KISEYRHYNR	<u>554</u>	58	140	17	161	1579	
ITIRCIICQR	<u>555</u>	488	93	50	123	12	
ILIRCIICQK	556	192	78	1383	1423	165	
WVGRCAACWR	557	2757	3973	360	24	19	
WAGRCAACWK	558	4662	583	23311	1491	50	
CFACWRSRRR	559	23542	7164	578	165	10206	
DTSIACVYCK	560	2936	89	5385	1968	216	
DVSIACVYCR	561	2814	217	406	487	658	
CVYCKATLEK	562	418	653	5307	17928	862	
RFEVYQFAFK	563	38	611	179	2867	2443	
RTEVYQFAFR	564	217	78	12	142	147	
AVKDLCIVYR	565	841	66	7.3	8.0	6.5	
AFKDLCIVYK	566	856	47	39	263	378	
ATCHKCIDFY	567	133	7.4	1164	12691	1386	
AACHKCIDFK	568	118	20	437	53733	414	
NLVYGETLEK	569	846	143	761	121	87	
NSVYGETLER	570	150	25	163	1333	18	
LSIRCLRCQK	571	245	14	100	1135	17	
LLIRCLRCOY	572	727	452	2894	2430	254	
RVHSIAGQYR	572 573	31	34	7.6	812	28	
RFHSIAGQYK	<u>575</u> 574	17	43	1.3	629	83	
LVTDLRIVYR	<u>574</u> 575	3869	648	20	150	6.8	
LFTDLRIVYK	<u>373</u> 576	628	263	258	149	277	
CTMCLRFLSK	<u>576</u> 577	1002	226	6274	3945	429	
CIMCLRFLSR	578	41	101	167	83	155	

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	SEQ ID						
Sequence	NO.	A*0301	A*1101	A*3101	A*3301	A*6801	
RLLSKISEYR	579	5.2	662	7.7	108	21	
RFLSKISEYY	<u>580</u>	1702	25535	14	41096	3999	
SFYGKTLEER	<u>581</u>	642	205	17	66	42	
SLYGKTLEEK	582	7.9	6.8	1044	6516	29	
WFGRCSECWR	583	1788	1569	20	5.5	26	
WTGRCSECWK	<u>584</u>	2492	26	3323	720	22	
AFCRVCLLFY	585	509	272	1777	1202	173	
AVCRVCLLFR	586	20	1.8	2.1	64	21	
CFLFYSKVRK	587	125	96	81	315	172	
CLLFYSKVRR	588	417	204	159	386	242	
LVYSKVRKYR	589	320	619	17	49	31	
LFYSKVRKYK	590	680	2582	18	30	1976	
GTTLESITKK	591	622	108	85182	132509	10147	
WVGSCLGCWR	592	48682	5520	20	15	9.3	
WTGSCLGCWK	593	7705	6.9	18344	2980	3.7	
VVADLRIVYR	594	513	18	41	101	16	
VFADLRIVYK	595	2086	127	402	200	273	
RTLSKISEYR	596	77	100	52	189	133	
RLLSKISEYK	597	15	65	158	40019	429	
KVSEYRHYNY	598	349	110	1791	70859	3498	
KISEYRHYNK	599	29	18	397	24827	15565	
IVIRCIICQR	600	984	217	52	529	28	
WLGRCAVCWR	601	2330	3002	356	40	112	
WTGRCAVCWK	602	1261	131	4176	3403	29	
YVVCDKCLK	603	3282	643	8.5	165	1289	
YAVCDKCLR	604	458	194	4261	26582	16034	
SVCRSSRTR	605	323	97	249	547	17	
SCCRSSRTK	606	21	3.9	51	5227	4.2	
SLPHAACHK	607	32	66	219	1186	654	
SIPHAACHR	608	1053	352	236	253	181	
FVDLTIVYR	609	29674	5312	2384	430	138	
FTDLTIVYK	610	557	16	24170	18477	143	
SFYGTTLEK	611	34	15	517	3385	498	
SVYGTTLER	612	28	6.4	133	454	21	
TFLEKLTNK	613	6839	815	451	148	918	
TTLEKLTNR	614	1993	817	42	37	101	
ETNPFGICK	615	9585	100	29103	804	14	
EGNPFGICR	616	11467	10372	5123	344	82	
NTLEOTVKR	617	20380	1151	2273	18	8.6	
ALCWRSRRR	618	959	9748	72	1289	7416	
AACWRSRRK	619	75	770	3022	45341	12877	
VSIACVYCR	620	3236	143	42	1347	185	
SIACVYCKK	621	271	83	9114	19632	96	
ILYRDCIAY	622	261	1832	53232	44670	>19607.84	
IVYRDCIAR	623	465	106	27	325	64	
CTAYAACHK	624	726	196	2956	771	167	
CIAYAACHR	625	3625	1905	502	115	262	
SFYGETLEK	626	288	108	947	885	1074	
	020						

		HLA-A3	HLA-A3 SUPERTYPE							
Sequence	SEQ ID NO.	A*0301	A*1101	A*3101	A*3301	A*6801				
SVYGETLER	627	44	11	235	160	17				
LIRCLRCQR	628	21335	12648	695	810	200				
RTQCVQCKK	629	234	20	127	8147	3066				
RLQCVQCKR	630	2535	6081	65	1829	11479				
KFLEERVKK	631	5344	2229	30	9740	17674				
KTLEERVKR	632	1957	159	37	1360	17685				
NVMGRWTGR	633	3884	794	40	18	20				
NIMGRWTGK	634	52	54	3274	86	173				
LTYRDDFPY	635	8265	82	>71146.25	20186	1529				
LVYRDDFPK		317	13	3009	1970	130				
RFCLLFYSK	<u>636</u>	1156	484	83	450	232				
RVCLLFYSR	637	439	111	51	2176	689				
	<u>638</u>	. 3.8	8.0	87	3382	13				
LTFYSKVRK	<u>639</u>	. 3.8 56	73	38	276	11				
LLFYSKVRR	<u>640</u>									
ATLESITKR	<u>641</u>	1437	16	100	851	188				
KVLCDLLIR	<u>642</u>	363	169	66	5896	9053				
KQLCDLLIK	<u>643</u>	226	65	340	46426	11897				
TFVHEIELK	<u>644</u>	4431	217	8412	4130	172				
TSVHEIELR	<u>645</u>	>64327.49	872	1039	5948	12				
YTFVFADLR	<u>646</u>	3633	8.1	20	6.6	2.9				
DFLEQTLKK	<u>647</u>	>57591.62	18809	34365	174	14376				
DTLEQTLKR	<u>648</u>	31347	12909	38127	9.2	110				
LVRCIICQR	<u>649</u>	677	358	59	109	201				
LIRCIICQK	<u>650</u>	445	252	639	834	285				
RVAVCWRPR	<u>651</u>	5.3	8.5	7.0	102	33				
RCAVCWRPK	<u>652</u>	285	340	382	131	1297				
AFCWRPRRR	<u>653</u>	273	17907	60	75	1087				
AVCWRPRRK	654	34	101	263	7950	1810				
LSFVCPWCA	655	38337	10864	4289	4603	341				
TFCCKCDSTLR	656	21772	8043	332	91	260				
LVVESSADDLR	657	>47008.55	2170	26410	5624	28				
TLQVVCPGCAR	658	20997	1395	67	63	147				
YLIHVPCCECK	659	1748	1534	33044	8066	177				
FVVQLDIQSTK	660	3682	853	48593	31350	2.7				
HTCNTTVR	661	4862	1792	726	4490	25				
GLVCPICSQK	662	428	814	45293	70317	3568				
GFNHQHLPAR	663	>46610.17	27889	173	5572	34617				
GVNHOHLPAK	664	42	11	3337	76239	9347				
NVVTFCCOCK	665	790	303	4757	87	13				
NIVTFCCQCR	666	1507	1070	2731	766	93				
GVSHAQLPAK	667	42	12	36011	>74935.4	20590				
LIHVPCCECR	668	5326	5925	385	387	228				
AVLQDIVLH	669	1922	101	6307	25776	27035				
ATLQDIVLK	670	37	8.6	65	17121	3231				
GVNHOHLPK	671	26	7.7	353	15615	1192				
HVMLCMCCK	672	282	7.7	772	825	99				
HTMLCMCCR	672 673	405	92	11	14	24				
LSFVCPWCR	673 674	31676	200	47	231	152				
LDI VCF WCK	0/4	31070	200	47	231	132				

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Sequence	SEQ ID NO.	A*0301	A*1101	A*3101	A*3301	A*6801	
AOPATADYK	675	3500	109	10413	58871	24173	
VVHAQLPAR	676	423	127	3.4	12	201	
VSHAQLPAK	677	378	9.5	46	1401	13502	
QLARQAKQH	678	8423	6862	945	1665	243	
KOHTCYLIR	679	135	213	13	2275	12177	
VTLDIQSTK	680	78	13	2046	1954	237	
VQLDIQSTR	681	15105	2917	162	4588	10341	
SLGPGPGTK	682	7.8	5.8	4392	152133	3517	
SLFGPGPGK	683	3.4	2.3	1085	82275	36	
LVGPGPGK	684	1004	291	23907	>125541.13	598	
KMFLOLAK	685	45	62	677	>125541.13	8384	
KMGPGPGK	686	84	242	1144	106362	4156	
KOENWYSLKK	687	608	178	6327	>136150.23	4794	
GVGPGPGLK	688	47	4.0	1367	>111538.46	3972	
GVSGPGPGK	689	13	5.8	>11221.95	>111538.46	209	
FLLYILFLVK	690	446	1431	54496	3254	2266	
LVFSNVLCFR	691	120	19	33	19	7.7	
SSFDIKSEVK	692	1900	19	19829	70344	31	
TLYOIOVMKR	693	361	164	397	558	90	
KOVOMMIMIK	694	264	112	4627	1231	2247	
GVIYIMIISK	695	777	18	18811	1567	1134	
ELFDKDTFFK	696	144	109	3676	13	3.6	
ALERLLSLKK	697	147	822	33559	18255	22391	
KILIKIPVTK	698	13	60	1661	24992	19571	
RLPLLPKTWK	699	11	67	340	11392	2889	
SOVSNSDSYK	700	1656	83	24559	>17448.86	1384	
QQNQESKIMK	701	3469	77	28120	>17448.86	21310	
HALLIPPK	702	30	5.3	23822	8426	82	
SSPLFNNFYK	703	100	0.7	1608	1728	6.3	
FLYLLNKKNK	704	177	475	4313	780	155	
LOMMNVNLOK	705	25	7.2	435	1113	320	
LTNHLINTPK	706	11	5.9	62	373	10	
IFISFYLINK	707	1987	1056	462	394	363	
RLFEESLGIR	707	64	1096	297	788	409	
LLYILFLVK	709	13	207	90687	13261	5545	
KSMLKELIK	710	189	151	450	>46548.96	>37037.04	
PVLTSLFNK	711	1949	25	5107	18271	29928	
KTMNNYMIK	712	17	5.5	24	12743	29	
LFDKDTFFK	713	931	167	5706	1189	101	
YLFNOHIKK	714	14	7.8	4919	7974	14	
MOSSFFMNR	715	13	1.1	29	75	3.8	
RFYITTRYK	716	1.9	67	15	98	17468	
TTRYKYLNK	710 717	117	848	416	652	2565	
AVIFTPIYY	717	25	9.5	42321	10068	1352	
ALERLLSLK	718 719	233	369	3433	12786	13708	
SISGKYDIK	720	2086	50	28249	12437	1745	
EORLPLLPK	721	1088	765	423	987	1911	
IALLIIPPK	722	1241	108	2926	1404	1965	

	HLA-A3 SUPERTYPE							
Sequence	SEQ ID NO.	A*0301	A*1101	A*3101	A*3301	A*6801		
PVVCSMEYK	723	1940	80	330791	22608	414		
VVCSMEYKK	724	443	54	891	14328	167		
FSYDLRLNK	725	29	4.9	461	1264	15		
HLNIPIGFK	726	2.3	1.3	183	97	2.8		
PLFNNFYKR	727	2635	1890	520	1258	132		
YQNFQNADK	728	2712	177	44698	>18447.84	19830		
QMMNVNLQK	729	20	7.0	504	6649	243		
AVSEIQNNK	730	25	11	1429	25449	14		
GTMYILLKK	731	2.2	1.2	29	8453	3.1		
FISFYLINK	732	19	9.0	2192	1456	18		
YLINKHWQR	733	1034	676	4.4	7.7	3.7		
ALKISQLQK	734	15	96	3203	23800	>54794.52		
KINSNFLLK	735	17	6.4	68	47740	2737		
AAMXDPTTFK	736	50	7.2					
GTMTTSXYK	737	4.0	4.5					
SXXPAXFQK	738	14	2.0					
ATAGDGXXEXRK	739	184	19					

TABLE 17

Sequence	SEO	AA	A-A24 SUP Organism		Position	Analog
Sequence	ID`	74.74	Ç. B3III			
AYGPGPGKF	NO.	9	Artificial	Consensus		A
	<u>740</u>		sequence			
AYIGPGPGF	741	9	Artificial	Consensus		Α
AYAAAAAAL	741	9	sequence Artificial			Poly
	<u>742</u>		sequence			•
AYSSWMYSY	<u>743</u>	9	EBV	EBNA3	176	
DLLDTASALY	<u>744</u>	10	HBV	Core	419	
WFHISCLTF	<u>745</u>	9	HBV	NUC	102	
KYTSFPWL	<u>746</u>	8	HBV	pol	745	
FAAPFTQCGY	<u>747</u>	10	HBV	pol	631	
SYQHFRKLLL	748	10	HBV	POL	4	
LYSHPIILGF	<u>749</u>	10	HBV	POL	492	
MSTTDLEAY	<u>750</u>	9	HBV	x	103	
MYVGDLCGSVF	<u>751</u>	1 I	HCV	EI	275	
MYGPGPGGSVF	<u>752</u>	11	HCV	EI	275	Α
MYVGPGPGSVF	<u>753</u>	11	HCV	El	275	Α
MYVGGPGPGVF	<u>754</u>	11	HCV	EI	275	Α
MYVGDGPGPGF	<u>755</u>	11	HCV	EI	275	Α
VMGSSYGF	<u>756</u>	8	HCV	NS5	2639	
EVDGVRLHRY	<u>757</u>	10	HCV	NS5	2129	
KYSKSSIVGW	<u>758</u>	10	HIV	NEF	4	Α
KWSKSSIVGF	<u>759</u>	10	HIV	NEF	4	Α
FFLKEKGGF	<u>760</u>	9	HIV	NEF	116	Α
IYSKKRQEF	<u>761</u>	9	HIV	NEF	175	Α
IYSKKRQEIF	<u>762</u>	10	HIV	NEF	175	Α
LYVYHTQGYF	<u>763</u>	10	HIV	NEF	190	Α
VYHTQGYFPDF	<u>764</u>	ΙI	HIV	NEF	192	Α
RYPLTFGW	765	8	HIV	NEF	216	
RYPLTFGF	<u>766</u>	8	HIV	NEF	216	Α
RFPLTFGF	<u>767</u>	8	HIV	NEF	216	Α
TYGWCFKL	<u>768</u>	8	HIV	NEF	222	Α
TFGWCFKF	<u>769</u>	8	HIV	NEF	222	Α
LYVYHTQGY	<u>770</u>	9	HIV	NEF	190	Α
NYTPGPGIRF	<u>771</u>	10	HIV	NEF	206	Α
QYPPLERLTL	772	10	HIV	REV	78	Α
QLPPLERLTF	<u>773</u>	10	HIV	REV	78	Α
KYGSLQYLAL	<u>774</u>	10	HIV	VIF	146	Α
LSKISEYRHY	<u>775</u>	10	HPV	E6	70	
ISEYRHYNY	<u>776</u>	9	HPV	E6	73	
RFHNIRGRW	777	9	HPV	E6	131	
RFLSKISEY	778	9	HPV	E6	68	
RFHNISGRW	779	9	HPV	E6	124	
VYDFAFRDLCI	780	H	HPV	E6	49	
PYAVCDKCLKF	781	11	HPV	E6	66	
QYNKPLCDLLI	782	H	HPV	E6	98	

Sequence	SEQ	AA		UPERTYPE m Protein	Position	Angl
sequence	ID NO.	AA	Organis	m riotem	rosition	Andi
PFGICKLCLRF	783	11	HPV	E6	59	
VYQFAFKDLCI	784	11	HPV	E6	44	
AYAACHKCIDF	785	11	HPV	E6	61	
VYKFLFTDLRI	786	11	HPV	E6	42	
PYGVCIMCLRF	787	11	HPV	E6	59	
PYAVCRVCLLF	788	11	HPV	E6	62	
VYDFVFADLRI	789	11	HPV	E6	42	
QYNKPLCDLF	790	10	HPV	E6	98	Α
VYEFAFKDLF	791	10	HPV	E6	44	Α
FYSKVSEFRF	792	10	HPV	E6	69	Α
VYREGNPFGF	793	10	HPV	E6	53	Α
FYSRIRELRF	794	10	HPV	E6	71	Α
PYAVCRVCLF	795	10	HPV	E6	62	Α
FYSKVRKYRF	796	10	HPV	E6	72	Α
LYGDTLEQTF	797	10	HPV	E6	83	Α
VYDFAFRDF	798	9	HPV	E6	49	Α
AYRDLCIVY	799	9	HPV	E6	53	Α
AFRDLCIVF	800	9	HPV	E6	53	Α
PYAVCDKCF	801	9	HPV	E6	66	Α
KYYSKISEY	802	9	HPV	E6	75	Α
KFYSKISEF	803	9	HPV	E6	75	Α
CYSLYGTTF	804	9	HPV	E6	87	Α
RYHNIRGRW	805	9	HPV	E6	131	Α
RFHNIRGRF	806	9	HPV	E6	131	Α
VYCKTVLEF	807	9	HPV	E6	33	Α
AYKDLFVVY	808	9	HPV	E6	48	Α
AFKDLFVVF	809	9	HPV	E6	48	Α
LYVVYRDSI	810	9	HPV	E6	52	Α
LFVVYRDSF	811	9	HPV	E6	52	Α
RYHNIAGHY	812	9	HPV	E6	126	Α
RFHNIAGHF	813	9	HPV	E6	126	Α
VYGTTLEKF	814	9	HPV	E6	83	Α
AYADLTVVY	815	9	HPV	E6	46	Α
AFADLTVVF	816	9	HPV	E6	46	Α
RYLSKISEY	817	9	HPV	E6	68	Α
NYSVYGNTF	818	9	HPV	E6	80	Α
RYHNISGRW	819	9	HPV	E6	124	Α
AYKDLCIVY	820	9	HPV	E6	48	Α
AFKDLCIVF	821	9	HPV	E6	48	Α
AYAACHKCF	822	9	HPV	E6	61	Α
VYGETLEKF	823	9	HPV	E6	85	Α
RYHSIAGQY	824	9	HPV	E6	126	Α
RFHSIAGQF	825	9	HPV	E6	126	Α
KYLFTDLRI	826	9	HPV	E6	44	Α
KFLFTDLRF	827	9	HPV	E6	44	Α
LYTDLRIVY	828	9	HPV	E6	46	Α
LFTDLRIVF	829	9	HPV	E6	46	Α

HLA-A24 SUPERTYPE								
Sequence	SEQ	AA	Organism	Protein	Position	Analog		
	ID NO.							
PYGVCIMCF	830	9	HPV	E6	59	Α		
RFLSKISEF	831	9	HPV	E6	68	Α		
EYRHYQYSF	832	9	HPV	E6	75	Α		
RYHNIMGRW	833	9	HPV	E6	124	Α		
RFHNIMGRF	834	9	HPV	E6	124	Α		
VYNFACTEF	835	9	HPV	E6	45	Α		
NYACTELKL	836	9	HPV	E6	47	Α		
NFACTELKF	837	9	HPV	E6	47	Α		
PYAVCRVCF	838	9	HPV	E6	62	Α		
LYYSKVRKY	839	9	HPV	E6	71	Α		
LFYSKVRKF		9	HPV	E6	71	A		
VYDFVFADF	840	9	HPV	E6	42	A		
VYADLRIVY	841	9	HPV	E6	46	Α		
VFADLRIVF	842	9	HPV	E6	46	A		
NYSLYGDTF	843	9	HPV	E6	80	A		
RFHNISGRF	844	9	HPV	E6	124	Ā		
LYNLLIRCF	845	9	HPV	E6	98	Ā		
FYSKVSEF	<u>846</u>	8	HPV	E6	69	^		
	847		HPV	E6	53			
VYREGNPF	<u>848</u>	8			44			
VFEFAFKDLF	<u>849</u>	10	HPV	E6	82			
EYRHYCYSLY	<u>850</u>	10	HPV	E6	75			
EYRHYNYSLY	<u>851</u>	10	HPV	E6				
ETRHYCYSLY	<u>852</u>	10	HPV	E6	82	A		
EYDHYCYSLY	<u>853</u>	10	HPV	E6	82	A		
KTRYYDYSVY	<u>854</u>	10	HPV	E6	78	A		
KYDYYDYSVY	<u>855</u>	10	HPV	E6	78	A		
ETRHYNYSLY	<u>856</u>	10	HPV	E6	75	A		
EYDHYNYSLY	<u>857</u>	10	HPV	E6	75	A		
TYCCKCDSTL	<u>858</u>	10	HPV	E7	56	A		
TFCCKCDSTF	<u>859</u>	10	HPV	E7	56	Α		
TYCHSCDSTF	<u>860</u>	10	HPV	E7	58	Α		
CYTCGTTVRF	<u>861</u>	10	HPV	E7	59	Α		
LYPEPTDLF	<u>862</u>	9	HPV	E7	15	Α		
NYYIVTCCF	863	9	HPV	E7	52	Α		
LFLNTLSF	864	8	HPV	E7	89			
LFLSTLSF	865	8	HPV	E7	90			
RVLPPNWKY	866	9	Human	40s ribo prot S13	132			
RLAHEVGWKY	867	10	Human	60s ribo prot L13A	139			
AYKKQFSQY	868	9	Human	60s ribo prot L5	217			
KTKDIVNGL	869	9	Human	Factin capping protein	235			
SLFVSNHAY	870	9	Human	fructose biphosphatealdolase	355			
TYGPGPGSLSF	871	11	Human	Her2/neu	63	Α		
TYLGPGPGLSF	872	11	Human	Her2/neu	63	Α		
TYLPGPGPGSF	873	11	Human	Her2/neu	63	Α		
TYLPTGPGPGF	874	11	Human	Her2/neu	63	Α		
RWGLLLALL	875	9	Human	Her2/neu	8			

HLA-A24 SUPERTYPE								
Sequence	SEQ	AA	Organism	Protein	Position	Analog		
	ID NO.							
PYVSRLLGI	876	9	Human	Her2/neu	780			
TYLPTNASL	877	9	Human	Her2/neu	63			
IYGPGPGLIF	878	10	Human	MAGE3	195	Α		
IYPGPGPGIF	879	10	Human	MAGE3	195	Α		
IYPKGPGPGF	880	10	Human	MAGE3	195	Α		
RISGVDRYY	330	9	Human	NADH	53			
	<u>881</u>			ubiqoxidoreductase				
LYSACFWWL	<u>882</u>	9	Human	OAI	194			
LYSACFWWF	883	9	Human	OAI	194	Α		
TYSVSFDSLF	<u>884</u>	10	Human	PSM	624			
TYGPGPGSLF	885	10	Human	PSM	624	Α		
TYSGPGPGLF	886	10	Human	PSM	624	Α		
TYSVGPGPGF	887	10	Human	PSM	624	Α		
AYPNVSAKI	888	9	Lysteria	listeriolysin	196			
AYGPGPGKI	889	9	Lysteria	listeriolysin	196	Α		
IMVLSFLF	890	8	Pf	CSP	427			
YYGKQENW	891	8	Pf	CSP	55			
VFNVVNSSI	892	9	Pf	CSP	416			
ALFQEYQCY	893	9	Pf	CSP	18			
LYNTEKGRHPF	894	11	Pf	EXP	100			
YFILVNLL	895	8	Pf	LSA	10			
KFFDKDKEL	896	9	Pf	LSA	76			
KFIKSLFHI	897	9	Pf	LSA	1876			
YFILVNLLIF	898	10	Pf	LSA	10			
FYFILVNLLIF	899	11	Pf	LSA	9			
SFYFILVNLLI	900	11	Pf	LSA	8			
VFLIFFDLF	901	9	Pf	SSP2	13			
LYLLMDCSGSI	902	11	Pf	SSP2	49			
KVSDEIWNY	903	9	Pf		182			
SYKSSKRDKF	904	10	Pf		225			
RYQDPQNYEL	905	10	Pf		21			
DFFLKSKFNI	906	10	Pf		3			
IFHFFLFLL	907	9	Pf		11			
VFLVFSNVL	908	9	Pf		41			
TYGIIVPVL	909	9	Pf		160			
NYMKIMNHL	910	9	Pf		34			
TYKKKNNHI	911	9	Pf		264			
VYYNILIVL	912	9	Pf		277			
LYYLFNOHI	913	9	Pf		285			
SFFMNRFYI	914	9	Pf		310			
FYITTRYKY	915	9	Pf		316			
KYINFINFI	915 916	9	Pf		328			
KYEALIKLL		9	Pf		380			
IYYFDGNSW	917	9	Pf		40			
VYRHCEYIL	918 010	9	Pf		94			
TWKPTIFLL	919	9	Pf		135			
SYKVNCINF	920	9	Pf		168			
0	<u>921</u>	-	• •					

Seguence	SEO	A A	Organism	Sequence SEQ AA Organism Protein Position Analog								
Sequence	ID	/4/1	O i Ballisiii		. 53111011							
	NO.											
KYNYFIHFF	922	9	Pf		216							
NYFIHFFTW	923	9	Pf		218							
HFFTWGTMF	924	9	Pf		222							
MFVPKYFEL	925	9	Pf		229							
IYTIIQDQL	926	9	Pf		295							
FFLKSKFNI	927	9	Pf		4							
RMTSLKNEL	928	9	Pf		61							
YYNNFNNNY	929	9	Pf		77							
YYNKSTEKL	930	9	Pf		87							
EYEPTANLL	931	9	Pf		109							
VYXKHPVSX	932	9	Unknown	Naturally processed		Α						
TYGNXTVTV	933	9	Unknown	Naturally processed		Α						
KYPDRVVPX	934	9	Unknown	Naturally processed		Α						
VYVXSXVTX	935	9	Unknown	Naturally processed		Α						
DAQXXXNTX	936	9	Unknown	Naturally processed		Α						
KYQAVTTTL	937	9	Unknown	Tumor p198	197							
KYGPGPGTTTL	938	11	Unknown	Tumor p198	197	Α						
KYOGPGPGTTL	939	11	Unknown	Tumor p198	197	Α						

TABLE 18

HLA-A24 SUPERTYPE								
	SEQ							
	ID							
Sequence	NO.	A*2402	A*2301	A*2902	A*3002			
AYGPGPGKF	<u>740</u>	2.4	9.7	44854	3.2			
AYIGPGPGF	<u>741</u>	217	12	15887	5728			
AYAAAAAAL	<u>742</u>	443						
AYSSWMYSY	<u>743</u>		21		4.9			
DLLDTASALY	744			74	37			
WFHISCLTF	745	204	11	95	75094			
KYTSFPWL	<u>746</u>	208	177	>172413.79	346			
FAAPFTQCGY	<u>747</u>			461	1364			
SYQHFRKLLL	<u>748</u>	418	39	28	3768			
LYSHPIILGF	<u>749</u>	2.6	5.4	109	1116			
MSTTDLEAY	<u>750</u>			2565	396			
MYVGDLCGSVF	<u>751</u>	26	0.91	612	1460			
MYGPGPGGSVF	<u>752</u>	35	5.4	48442	31980			
MYVGPGPGSVF	<u>753</u>	35	4.4	1527	28177			
MYVGGPGPGVF	<u>754</u>	381	85	89	2870			
MYVGDGPGPGF	755	90	11	8656	39608			
VMGSSYGF	<u>756</u>	36	159	145	41967			
EVDGVRLHRY	757			14940	113			
KYSKSSIVGW	758	4061	491	>69444.44	>34482.76			
KWSKSSIVGF	759	1674	84	>56179.78	30367			
FFLKEKGGF	760	3456	655	3015	141			
IYSKKROEF	761	306	421	29353	727			
IYSKKRQEIF	762	238	360	>131578.95	21001			
LYVYHTOGYF	763	38	23	1696	1222			
VYHTQGYFPDF	764	149	68	14923	>22556.39			
RYPLTFGW	765	127	3836	13889	6251			
RYPLTFGF	766	3.3	6.4	9704	6328			
RFPLTFGF	767	178	124	12759	13472			
TYGWCFKL	768	2181	333	25658	>8042.9			
TFGWCFKF	769	3424	462	4449	>10135.14			
LYVYHTOGY	770	7140	6088	216	258			
NYTPGPGIRF	771	483	37	8334	>9646.3			
QYPPLERLTL	772	211	22	>11520.74	>9646.3			
OLPPLERLTF	773	2507	338	>37313.43	>36585.37			
KYGSLQYLAL	774	2800	147	>69444.44	6957			
LSKISEYRHY	775	>93023.26	>23671.5	55190	186			
ISEYRHYNY	776	125794	>23557.69	1329	32			
RFHNIRGRW	777	53237	11416	18	58			
RFLSKISEY	778	472	121	34623	23			
RFHNISGRW	779	>80536.91	22871	174	37			
VYDFAFRDLCI	780	44	8.9	62242	35724			
PYAVCDKCLKF	781	99	8.1	118249	>60000			
OYNKPLCDLLI	782	303	36	>166666.67	6680			

HLA-A24 SUPERTYPE									
	SEQ								
	ID								
Sequence	NO.	A*2402	A*2301	A*2902	A*3002				
PFGICKLCLRF	<u>783</u>	137	19	1249	32803				
VYQFAFKDLCI	<u>784</u>	30	1.9	49276	3477				
AYAACHKCIDF	<u>785</u>	91	14	1264	4699				
VYKFLFTDLRI	<u>786</u>	37	14	30216	1865				
PYGVCIMCLRF	<u>787</u>	380	100	69	43722				
PYAVCRVCLLF	788	226	150	2711	53351				
VYDFVFADLRI	789	47	8.0	8904	7585				
OYNKPLCDLF	790	115	21	7658	525				
VYEFAFKDLF	791	15	1.7	1973	2038				
FYSKVSEFRF	792	7.1	2.2	79	18453				
VYREGNPFGF	793	197	91	11120	21947				
FYSRIRELRF	794	11	1.6	83	12598				
PYAVCRVCLF	795	12	4.5	407	5226				
FYSKVRKYRF	796	18	13	3042	1232				
LYGDTLEOTF	797	91	24	40871	42025				
VYDFAFRDF	798	9.6	19	47381	8490				
AYRDLCIVY	799	2094	1479	7117	66				
AFRDLCIVF	800	1005	369	6722	3305				
PYAVCDKCF	801	216	183	122025	9884				
KYYSKISEY	802	10951	2165	702	1.3				
KFYSKISEF	803	174	138	73339	306				
CYSLYGTTF	804	28	11	2088	7823				
RYHNIRGRW	805	145	14	122644	15				
RFHNIRGRF	806	29	2.4	346	0.69				
VYCKTVLEF	807	50	4.7	610	1139				
AYKDLFVVY	808	1549	905	639	1.3				
AFKDLFVVF	809	294	6.8	3051	829				
LYVVYRDSI	810	982	242	148359	3483				
LFVVYRDSF	811	268	134	919	18				
RYHNIAGHY	812	1227	195	138	0.93				
RFHNIAGHF	813	37	17	635	1.4				
VYGTTLEKF	814	19	13	75267	220				
AYADLTVVY	815	369	1384	136	9.3				
AFADLTVVF	816	203	30	779	137				
RYLSKISEY	817	142	98	4247	1.1				
NYSVYGNTF	818	28	29	9121	2559				
RYHNISGRW	819	47	15	104884	13				
AYKDLCIVY	820	33798	3036	5205	29				
AFKDLCIVF	821	284	16	5846	2305				
AYAACHKCF	822	200	159	10972	3393				
VYGETLEKF	823	45	14	91902	20009				
RYHSIAGOY	824	3170	1904	544	1.4				
RFHSIAGOF	825	28	2.9	481	1.2				
KYLFTDLRI	826	108	1.9	78575	339				
KFLFTDLRF	827	12	0.74	44	152				
LYTDLRIVY	828	1986	1216	4.8	2.1				

	Н	LA-A24 SUPI	ERTYPE		
	SEQ				
	ID				
Sequence	NO.	A*2402	A*2301	A*2902	A*3002
LFTDLRIVF	829	169	2.6	164	2649
PYGVCIMCF	<u>830</u>	190	147	144402	38850
RFLSKISEF	<u>831</u>	58	2.5	40103	201
EYRHYQYSF	<u>832</u>	21	2.3	13707	430
RYHNIMGRW	<u>833</u>	29	12	106990	7.1
RFHNIMGRF	834	39	2.6	174	1.3
VYNFACTEF	<u>835</u>	14	2.1	774	784
NYACTELKL	<u>836</u>	1741	131	77844	49107
NFACTELKF	837	211	13	46	6826
PYAVCRVCF	<u>838</u>	429	257	5602	316
LYYSKVRKY	<u>839</u>	21942	2735	1452	28
LFYSKVRKF	840	2008	277	11172	632
VYDFVFADF	841	9.9	2.2	1230	3961
VYADLRIVY	842	28	122	8.2	8.3
VFADLRIVF	843	23	2.5	87	24062
NYSLYGDTF	<u>844</u>	6.4	142	20945	64
RFHNISGRF	<u>845</u>	34	5.5	572	2.8
LYNLLIRCF	<u>846</u>	47	15	17958	2255
FYSKVSEF	847	21	18	3774	66667
VYREGNPF	<u>848</u>	554	147	10001	65970
VFEFAFKDLF	<u>849</u>	400			
EYRHYCYSLY	<u>850</u>			198	3.7
EYRHYNYSLY	<u>851</u>			956	12
ETRHYCYSLY	<u>852</u>			755	10
EYDHYCYSLY	<u>853</u>			799	77
KTRYYDYSVY	<u>854</u>			87841	0.71
KYDYYDYSVY	<u>855</u>			5749	11
ETRHYNYSLY	<u>856</u>			5464	29
EYDHYNYSLY	<u>857</u>			777	93
TYCCKCDSTL	<u>858</u>	206	30	145803	16588
TFCCKCDSTF	<u>859</u>	25	14	501	1167
TYCHSCDSTF	<u>860</u>	14	2.9	5236	3580
CYTCGTTVRF	<u>861</u>	41	18	7744	38331
LYPEPTDLF	<u>862</u>	38	17	1150	30732
NYYIVTCCF	<u>863</u>	27	12	2675	8398
LFLNTLSF	<u>864</u>	587	104	1013	118217
LFLSTLSF	<u>865</u>	2283	160	1034	>75000
RVLPPNWKY	<u>866</u>		>49000		3.0
RLAHEVGWKY	<u>867</u>		4631		3.8
AYKKQFSQY	<u>868</u>		10669		5.3
KTKDIVNGL	<u>869</u>		>49000		164
SLFVSNHAY	<u>870</u>		30295		1.1
TYGPGPGSLSF	<u>871</u>	7.1	1.7	9853	47246
TYLGPGPGLSF	<u>872</u>	23	0.65	600	26889
TYLPGPGPGSF	<u>873</u>	8.8	2.2	56183	7275
TYLPTGPGPGF	<u>874</u>	39	8.6	56574	32985

HLA-A24 SUPERTYPE								
	SEQ							
	ID							
Sequence	NO.	A*2402	A*2301	A*2902	A*3002			
RWGLLLALL	875	106	100	61253	300			
PYVSRLLGI	876	11	18	200160	65465			
TYLPTNASL	877	141	7.8	106153	8244			
IYGPGPGLIF	878	7.4	8.0	58	6845			
IYPGPGPGIF	879	58	12	18659	17959			
IYPKGPGPGF	880	7.5	4.9	53603	61283			
RISGVDRYY	881		>49000		3.0			
LYSACFWWL	882	28						
LYSACFWWF	883	28						
TYSVSFDSLF	884	10	12	521	5218			
TYGPGPGSLF	885	3.9	8.7	7228	10871			
TYSGPGPGLF	886	50	92	7726	3461			
TYSVGPGPGF	887	332	340	120913	55200			
AYPNVSAKI	888	14	45	56905	4456			
AYGPGPGKI	889	36	169	>156250	5427			
IMVLSFLF	890	469	7.5	111	30000			
YYGKQENW	891	85	951	>50000	>30000			
VFNVVNSSI	892	403	35	24001	15737			
ALFQEYQCY	893			149	1032			
LYNTEKGRHPF	894	175	1947	>50000	>30000			
YFILVNLL	895	96	82	4050	30000			
KFFDKDKEL	896	269	>49000	>50000	3012			
KFIKSLFHI	897	4.1	2.0	>50000	3495			
YFILVNLLIF	898	577	12	764	3388			
FYFILVNLLIF	899	599	50	902	9826			
SFYFILVNLLI	900	229	35	3066	2096			
VFLIFFDLF	901	40	12	1510	13554			
LYLLMDCSGSI	902	154	10	5893	1469			
KVSDEIWNY	903	52169	>11980.44	230	1.9			
SYKSSKRDKF	904	256	797	12594	88			
RYODPONYEL	905	212	124	79717	189			
DFFLKSKFNI	906	1648	304	47714	491			
IFHFFLFLL	907	208	80	1405	837			
VFLVFSNVL	908	26	4.9	33675	37689			
TYGIIVPVL	909	248	20	30056	1519			
NYMKIMNHL	910	16	1.7	45443	110			
TYKKKNNHI	911	30	81	21642	162			
VYYNILIVL	912	265	52	>192307.69	1127			
LYYLFNOHI	913	33	1.4	20130	11035			
SFFMNRFYI	914	172	11	200	1022			
FYITTRYKY	915	350	11	9.6	7.5			
KYINFINFI	916	11	0.72	25475	55			
KYEALIKLL	917	2856	484	17296	16098			
IYYFDGNSW	918	80	6.1	3101	3025			
VYRHCEYIL	919	2200	64	117851	3326			
TWKPTIFLL	920	148	11	21155	306			

	HLA-A24 SUPERTYPE									
	SEQ									
Sequence	<u>ID</u> NO.	A*2402	A*2301	A*2902	A*3002					
SYKVNCINF	921	27	15	2535	572					
KYNYFIHFF	922	2.5	0.49	319	2.7					
NYFIHFFTW	923	9.3	1.3	9774	3020					
HFFTWGTMF	<u>924</u>	83	5.7	4.0	220					
MFVPKYFEL	925	266	11	2560	8560					
IYTIIQDQL	<u>926</u>	72	45	>37313.43	14124					
FFLKSKFNI	927	1434	49	43105	>83333.33					
RMTSLKNEL	928	12711	1807	40270	14					
YYNNFNNNY	<u>929</u>	817	126	19	34					
YYNKSTEKL	930	109	106	55636	21751					
EYEPTANLL	931	127	44	>37313.43	>26086.96					
VYXKHPVSX	932	4.3								
TYGNXTVTV	933	26								
KYPDRVVPX	934	224								
VYVXSXVTX	935	5.3								
DAQXXXNTX	936	5.9								
KYQAVTTTL	937	22	16	>156250	625					
KYGPGPGTTTL	938	103	130	9180	7056					
KYQGPGPGTTL	939	543	438	74453	5999					

TABLE 19

			HLA-B7 SUP	ERTYPE		
	SEQ					
Sequence	ID NO.	AA		Protein	Position	
APGPGPGLL		9	Artificial	Consensus		Α
. nn ananat	<u>940</u>	9	sequence	Consensus		Α
APRGPGPGL	941	9	Artificial sequence	Consensus		А
OPRAPIRPI	942	9	EBNA		881	
YPLHEQHGM	942	9	EBNA		458	
CPTVQASKL	943	9	HBV	NUC	14	
SPTYKAFL	945	8	HBV	pol	659	
SPGPGPGL	945	8	HBV	pol	659	Α
TPAGPGPGVF		10	HBV	pol	354	A
TPARGPGPGF	947	10	HBV	pol	354	A
TPTGWGLAI	948	9	HBV	POL	691	
APCNFFTSA	949	9	HBV	X	146	
GPGHKARVI	950	9	HIV	GAG	390	Α
RPOVPLRPMTI	<u>951</u>	11	HIV	NEF	98	A
FPVRPQVPI	<u>952</u>	9	HIV	NEF	94	A
-	953	9	HIV	NEF	98	Â
RPQVPLRPI	<u>954</u>	11	HIV	NEF	98	A
RPQVPLRPMTI	<u>955</u>			NEF	217	A
YPLTFGWCI	<u>956</u>	9	HIV	NEF	217	A
FPLTFGWCI	<u>957</u>		HIV		217	A
FPLTFGWCFKI	958	11	HIV	NEF	94	А
FPVRPQVPL	<u>959</u>	9	HIV	nef	94	
FPGPGPGPL	<u>960</u>	9	HIV	nef		A
FPVGPGPGL	<u>961</u>	9	HIV	nef	94	A
GPKVKQWPI	<u>962</u>	9	HIV	POL	197	A
LPPLERLTI	963	9	HIV	REV	79	Α
CPEEKQRHL	<u>964</u>	9	HPV	E6	118	
VPGPGPGL	<u>965</u>	8	Human	Her2/neu	884	Α
RPGPGPGVSEF	<u>966</u>	11	Human	Her2/neu	966	Α
RPRGPGPGSEF	<u>967</u>	11	Human	Her2/neu	966	Α
RPRFGPGPGEF	<u>968</u>	11	Human	Her2/neu	966	Α
RPRFRGPGPGF	<u>969</u>	11	Human	Her2/neu	966	Α
APGPGPGAAPA	970	11	Human	p53	76	Α
APAGPGPGAPA	<u>971</u>	11	Human	p53	76	Α
APAAGPGPGPA	972	11	Human	p53	76	Α
APAAPGPGPGA	973	11	Human	p53	76	Α
RPRGDNFAV	974	9	Pf	SSP2	305	
RPGPGPGAV	975	9	Pf	SSP2	305	Α
RPRGPGPGV	976	9	Pf	SSP2	305	Α
APRTVALTAL	977	10	Unknown	Naturally procesed		
APGPGPGTAL	978	10	Unknown	Naturally procesed		Α
APRGPGPGAL	979	10	Unknown	Naturally procesed		Α
APRTGPGPGL	980	10	Unknown	Naturally procesed		Α
XVXDNATEY	981	9	Unknown	Naturally procesed		Α
LGFVFTLTV	982	9	unknown			

TABLE 20

HLA-B7 SUPERTYPE										
	SEO									
Sequence		B*0702	B*3501	B*5101	B*5301	B*5401				
APGPGPGLL	940	299	7481	1614	18117	15613				
APRGPGPGL	941	4.9	974	633	19779	1120				
QPRAPIRPI	942	6770	>72000	>55000	12	>100000				
YPLHEQHGM	943	>55000	20785	>55000	10	>100000				
CPTVQASKL	944	3247	645	448	1861	21643				
SPTYKAFL	945	109	31169	4665	54879	58651				
SPGPGPGL	946	173	2337	3535	25607	53272				
TPAGPGPGVF	947	334	374	296	2629	351				
TPARGPGPGF	948	144	1678	2418	2742	31768				
TPTGWGLAI	949	76	5145	103	1343	172				
APCNFFTSA	950	43	8087	1045	>22409.64	0.61				
GPGHKARVI	951	1686	>72000	>55000	2.2	>50000				
RPQVPLRPMTI	952	47009	>18997.36	8081	21518	129				
FPVRPQVPI	953	94	124	39	222	9.1				
RPQVPLRPI	954	367	>23225.81	>9001.64	85335	1215				
RPQVPLRPMTI	955	140	10455	5045	21538	>15128.59				
YPLTFGWCI	956	54283	1378	153	154	79				
FPLTFGWCI	957	47951	164	63	36	14				
FPLTFGWCFKI	958	52567	4991	590	188	105				
FPVRPQVPL	959	17	3.8	18	49	21				
FPGPGPGPL	960	1584	426	2330	21036	29900				
FPVGPGPGL	961	106	14	138	32	246				
GPKVKQWPI	962	5500	>72000	>55000	2.3	>50000				
LPPLERLTI	963	24398	13399	359	2624	11243				
CPEEKQRHL	964	10	>52554.74	>35483.87	>109411.76	>76923.08				
VPGPGPGL	965	1517	447	537	4094	46405				
RPGPGPGVSEF	966	119	18115	16774	20988	3360				
RPRGPGPGSEF	967	11	24871	>14824.8	19336	2745				
RPRFGPGPGEF	968	14	>30901.29	>14824.8	76844	15470				
RPRFRGPGPGF	969	9.7	>30901.29	>14824.8	49682	60095				
APGPGPGAAP		1112	1252	1317	4366	361				
A	<u>970</u>									
APAGPGPGAP		161	>28915.66	11947	>39743.59	43				
A A D A A CDCDCD	<u>971</u>	173	12845	12470	28574	204				
APAAGPGPGP A	972	1/3	12845	12470	28574	204				
APAAPGPGPG	212	811	3484	15814	>39240.51	158				
A	973									
RPRGDNFAV	974	12	20386	1681	>46268.66	212				
RPGPGPGAV	975	23	48487	2899	>46268.66	1891				
RPRGPGPGV	976	11	2368	52	34831	47				
APRTVALTAL	977	12	4351	14601	61596	16804				
APGPGPGTAL	978	81	16315	16462	>43661.97	35965				
APRGPGPGAL	979	11	23381	12732	>43661.97	1665				
APRTGPGPGL	980	15	1414	1559	22012	2043				
XVXDNATEY	981	>55000	444			>100000				
LGFVFTLTV	982	849	>72000	27500	>93000	464				

TABLE 21

		H	ILA-B44 SUP	ERTYPE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
SEAAYAKKI	110.	9	Artificial	pool consensus	1 03111011	Anarog
SEANTAKKI	983	,	sequence	poor conscisus		**
GEFPYKAAA	202	9	Artificial	pool consensus		Α
	984		sequence	•		
SEAPYKAIL		9	Artificial	pool consensus		Α
	<u>985</u>		sequence			
SEAPKYAIL		9	Artificial	pool consensus		Α
	<u>986</u>	•	sequence			
AEFKYIAAV	007	9	Artificial	pool consensus		Α
AEIPYLAKY	<u>987</u>	9	sequence Artificial	pool consensus		Α
AEIFILAKI	988	,	sequence	poor consensus		л
AEIPKLAYF	200	9	Artificial	pool consensus		Α
ALDII ILDII I	989	-	sequence	F		
FPFDYAAAF		9	Artificial			Α
	990		sequence			
FPFKYKAAF		9	Artificial			Α
	<u>991</u>		sequence			
FPFKYAKAF		9	Artificial			Α
PPPKKKAAAF	<u>992</u>		sequence			
FPFKYAAAF	993	9	Artificial sequence			Α
FAFKYAAAF	993	9	Artificial			Α
TATKTAAAT	994	,	sequence			**
FQFKYAAAF	22.	9	Artificial			Α
	<u>995</u>		sequence			
FDFKYAAAF		9	Artificial			Α
	<u>996</u>		sequence			
SENDRYRLL	<u>997</u>	9	EBV	BZLF1	209	A
IEDPPYNSL	<u>998</u>	9	EBV	lmp2	200	A
YEANGNLI	999	8	Flu	HA	259	A
YEDLRVLSF	<u>1000</u>	9	Flu	NP	338	Α
SDYEGRLI	1001	8	Flu	NP	50	
GEISPYPSL	<u>1002</u>	9	Flu	NS1	158	Α
MDIDPYKEF	<u>1003</u>	9	HBV	NUC	30	
LDKGIKPY	1004	8	HBV	POL	125	
ADLMGYIPL	1005	9	HCV	core	131	
LDPYARVAI	<u>1006</u>	9	HCV	NS5b	2663	Α
AENLWVTVY	1007	9	HIV	gp120	1	
KENLWVTVY	1008	9	HIV	gp120	1	A
AEKLWVTVY	1009	9	HIV	gp120	. 1	A
AENKWVTVY	<u>1010</u>	9	HIV	gp120	1	A
AENLKVTVY	1011	9	HIV	gp120	1	A
AENLWKTVY	1012	9	HIV	gp120	1	A A
AENLWVKVY	1013	9	HIV	gp120	1	A
AENLWVTKY AENLWVTVK	1014	9	HIV HIV	gp120 gp120	1	A
FENLWVTVY	1015	9	HIV		1	A
VENLWVTVY	1016	9	HIV	gp120 gp120	1	A
PENLWVTVY	1017	9	HIV	gp120 gp120	1	A
NENLWYTVY	1018	9	HIV	gp120 gp120	1	A
DENLWVTVY	1019 1020	9	HIV	gp120 gp120	1	A
DEMEN A I A I	1020	,	4 1 1 Y	Phiro	•	-

Sequence	SEQ ID					
Seguence						
	NO.	AA	Organism	Protein	Position	Analog
TENLWVTVY	1021	9	HIV	gp120	1	A
YENLWVTVY	1022	9	HIV	gp120	1	Α
ATNLWVTVY	1023	9	HIV	gp120	1	Α
AEFLWVTVY	1024	9	HIV	gp120	1	Α
AEVLWVTVY	1025	9	HIV	gp120	1	Α
AEPLWVTVY	1026	9	HIV	gp120	1	Α
AEDLWVTVY	1027	9	HIV	gp120	1	Α
AENLWVTVY	1028	9	HIV	gp120	1	
AETLWVTVY	1029	9	HIV	gp120	1	Α
AENFWVTVY	1030	9	HIV	gp120	1	Α
AENVWVTVY	1031	9	HIV	gp120	1	Α
AENPWVTVY	1032	9	HIV	gp120	1	Α
AENDWVTVY	1033	9	HIV	gp120	ī	A
AENNWVTVY	1034	9	HIV	gp120	1	A
AENTWVTVY	1035	9	HIV	gp120	i	A
AENLFVTVY	1036	9	HIV	gp120	1	A
AENLVVTVY	1037	9	HIV	gp120	i	A
AENLPVTVY	1037	9	HIV	gp120	i	A
AENLDVTVY	1039	9	HIV	gp120	1	A
AENLNVTVY	1040	9	HIV	gp120	1	A
AENLTVTVY	1040	9	HIV	gp120	i	A
AENLWFTVY	1041	9	HIV	gp120	1	A
AENLWLTVY	1042	9	HIV	gp120	i	A
AENLWPTVY	1043	9	HIV	gp120	1	A
AENLWDTVY	1044	9	HIV	gp120	i	A
AENLWNTVY	1045	9	HIV	gp120	i	A
AENLWTTVY	1048	9	HIV	gp120	i	A
AENLWVFVY	1047	9	HIV	gp120	i	A
AENLWVVVVY	1048	9	HIV	gp120	1	A
AENLWVPVY	1050	<u></u>	HIV	gp120	i	A
AENLWVDVY	1050	9	HIV	gp120	i	A
AENLWVNVY	1051	9	HIV	gp120	1	A
AENLWVSVY		9	HIV	gp120	1	A
AENLWVTFY	1053 1054	9	HIV	gp120	î	A
AENLWVTLY	1054	9	HIV	gp120	1	A
AENLWVTPY		9	HIV	gp120	1	A
AENLWVTDY	1056	9	HIV	gp120	1	A
AENLWVTNY	1057	9	HIV	gp120 gp120	1	A
AENLWVTTY	1058	9	HIV	gp120	1	A
AENLWVTVA	1059	9	HIV		1	A
AENLWVTVC	1060	9	HIV	gp120	1	A
	1061	9		gp120	1	A
AENLWVTVE AENLWVTVF	1062	9	HIV HIV	gp120	1	A
	1063			gp120	1	
AENLWVTVG	1064	9	HIV	gp120	1	A A
AENLWVTVH	1065		HIV	gp120	1	A
AENLWVTVI	1066	9	HIV	gp120	1	
AENLWVTVL	1067	9	HIV	gp120		A
AENLWVTVM	1068	9	HIV	gp120	1	A
AENLWVTVN	1069	9	HIV	gp120	1	A
AENLWVTVP	1070	9	HIV	gp120	1 1	A A
AENLWVTVQ	<u>1071</u>	9	HIV	gp120	1	A

		Н	ILA-B44 SUP	ERTYPE		
	SEQ					
Cognongo	ID NO.	AA	Organism	Protein	Position	Analog
Sequence AENLWVTVR	1072	9	HIV	gp120	1	Analog
AENLWVTVS	1072	é	HIV	gp120	i	A
AENLWVTVT	1074	9	HIV	gp120	i	A
AENLWVTVV	1075	9	HIV	gp120	i	A
AENLWVTVW	1075	9	HIV	gp120	1	A
AENLWVTVY	1076	9	HIV	gp120 gp120	1	А
AENLYVTVF	1077	9	HIV	gp120	1	Α
TEPAAVGVGAV	1078	11	HIV	NEF	33	А
AEPAAEGV	1079	8	HIV	NEF	34	
AEPAAEGVGA	_	10	HIV	NEF	34	
AEPAAEGVGAV	1081	11	HIV	NEF	34	
	1082	10		NEF	84	
QEEEEVGFPV	1083	9	HIV			
EEEEVGFPV	1084		HIV	NEF	86 87	
EEEVGFPV	1085	8	HIV	NEF		
EEVGFPVRPQV	1086	11	HIV	NEF	88	
DEEVGFPV	1087	8	HIV	NEF	89	
KEKGGLDGL	1088	9	HIV	NEF	120	
KEKGGLDGLI	1089	10	HIV	NEF	120	
QEILDLWV	<u>1090</u>	8	HIV	NEF	184	
QEILDLWVY	<u>1091</u>	9	HIV	NEF	184	
AETFYVDGA	<u>1092</u>	9	HIV	POL	629	
EEKPRTLHDL	<u>1093</u>	10	HPV	E6	6	
NEILIRCII	<u>1094</u>	9	HPV	E6	97	
QEKKRHVDL	<u>1095</u>	9	HPV	E6	113	
AEGKEVLL	<u>1096</u>	8	Human	CEA	46	
QELFIPNI	<u>1097</u>	8	Human	CEA	282	
QELFISNI	1098	8	Human	CEA	460	
TEKNSGLY	1099	8	Human	CEA	468	
AELPKPSI	1100	8	Human	CEA	498	
PEAQNTTY	1101	8	Human	CEA	525	
IESTPFNVA	1102	9	Human	CEA	38	
AEGKEVLLL	1103	9	Human	CEA	46	
EEATGQFRV	1104	9	Human	CEA	132	
VEDKDAVAF	1105	9	Human	CEA	157	
CEPETQDAT	1106	9	Human	CEA	167	
PETQDATYL	1107	9	Human	CEA	169	
CETQNPVSA	1108	9	Human	CEA	215	
QELFIPNIT	1109	9	Human	CEA	282	
AEPPKPFIT	1110	9	Human	CEA	320	
VEDEDAVAL	1111	9	Human	CEA	335	
CEPEIQNTT	1112	9	Human	CEA	345	
PEIQNTTYL	1113	9	Human	CEA	347	
YECGIQNEL	1114	9	Human	CEA	391	
QELFISNIT	1115	9	Human	CEA	460	
TEKNSGLYT	1116	9	Human	CEA	468	
AEGKEVLLLV	1117	10	Human	CEA	46	
KEVLLLVHNL	1118	10	Human	CEA	49	
GERVDGNRQI	1119	10	Human	CEA	70	
REIIYPNASL	1120	10	Human	CEA	98	
REIIYPNASL NEEATGOFRV	1120 1121	10 10	Human Human	CEA CEA	98 131	

		H	ILA-B44 SUP	ERTYPE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analo
GENLNLSCHA	1123	10	Human	CEA	252	
QELFIPNITV	1124	10	Human	CEA	282	
CEPEIQNTTY	1125	10	Human	CEA	345	
PEIQNTTYLW	1126	10	Human	CEA	347	
CEPEAQNTTY	1127	10	Human	CEA	523	
PEAQNTTYLW	1128	10	Human	CEA	525	
MESPSAPPHRW	1129	11	Human	CEA	1	
IESTPFNVAEG	1130	11	Human	CEA	38	
GERVDGNRQII	1131	11	Human	CEA	70	
REIIYPNASLL	1132	11	Human	CEA	98	
NEEATGOFRVY	1133	11	Human	CEA	131	
CEPETQDATYL	1134	11	Human	CEA	167	
GENLNLSCHAA	1134	11	Human	CEA	252	
CEPEIQNTTYL	1136	11	Human	CEA	345	
PEIQNTTYLWW		11	Human	CEA	347	
•	1137	11	Human	CEA	391	
YECGIQNELSV	1138	11	Human	CEA	397	
NELSVDHSDPV	1139				523	
CEPEAQNTTYL	1140	11	Human	CEA		
PEAQNTTYLWW	1141	11	Human	CEA	525	
PEIQNTTYLWWV	<u>1142</u>	12	Human	CEA	347	
PEAQNTTYLWW	1142	12	Human	CEA	525	
V CEREIONITEVI W	<u>1143</u>	13	Human	CEA	345	
CEPEIQNTTYLW W	1144	13	riuman	CLA	343	
W AEMGKGSFKY	1144	10	Human	elong. Factor Tu	48	
SEDCQSL	1146	7	Human	Her2/neu	209	
REVRAVT	1147	7	Human	Her2/neu	351	
FETLEEI	1148	7	Human	Her2/neu	400	
TELVEPL	1149	7	Human	Her2/neu	694	
SECRPRF	1150	7	Human	Her2/neu	963	
PETHLDML	1151	8	Human	Her2/neu	39	
QEVQGYVL	1152	8	Human	Her2/neu	78	
RELOLRSL	1152	8	Human	Her2/neu	138	
CELHCPAL	1154	8	Human	Her2/neu	264	
LEEITGYL		8	Human	Her2/neu	403	
EEITGYLY	1155 1156	8	Human	Her2/neu	404	
DECVGEGL	1156	8	Human	Her2/neu	502	
AEQRASPL	1157	8	Human	Her2/neu	644	
KEILDEAY	1158	8	Human	Her2/neu	765	
EEAPRSPL	_	8	Human	Her2/neu	1068	
SEDPTVPL	1160	8	Human	Her2/neu	1113	
MELAALCRW	1161	9	Human	Her2/neu	1	
	1162	9	Human	Her2/neu	78	
QEVQGYVLI	1163	9	Human	Her2/neu	108	
FEDNYALAV	1164	9	Human	Her2/neu	138	
RELQLRSLT	1165	9		Her2/neu	146	
TEILKGGVL	1166	9	Human		237	
HEQCAAGCT	1167		Human	Her2/neu		
CELHCPALV	1168	9	Human	Her2/neu	264	
FESMPNPEG	1169	9	Human	Her2/neu	279	
QEVTAEDGT	1170	9	Human	Her2/neu	320	
CEKCSKPCA	1171	-	Human	Her2/neu	331 347	
MEHLREVRA	1172	9	Human	Her2/neu	347	

	oric	Н	ILA-B44 SUP	EKIYPE		
	SEQ ID					
Sequence	NO.	AA	Organism	Protein	Position	Analo
REVRAVTSA	1173	9	Human	Her2/neu	351	
OEFAGCKKI	1174	9	Human	Her2/neu	362	
EEITGYLYI	1175	9	Human	Her2/neu	404	
RELGSGLAL	1176	9	Human	Her2/neu	459	
GEGLACHQL	1177	9	Human	Her2/neu	506	
QECVEECRV	1178	9	Human	Her2/neu	538	
VEECRVLOG	1179	9	Human	Her2/neu	541	
EECRVLQGL	1180	9	Human	Her2/neu	542	
AEQRASPLT	1181	9	Human	Her2/neu	644	
QETELVEPL	1182	9	Human	Her2/neu	692	
VEPLTPSGA	1183	9	Human	Her2/neu	697	
TELRKVKVL	1184	9	Human	Her2/neu	718	
GENVKIPVA	1185	9	Human	Her2/neu	743	
KEILDEAYV	1186	9	Human	Her2/neu	765	
DEAYVMAGV	1187	9	Human	Her2/neu	769	
DETEYHADG	1188	9	Human	Her2/neu	873	
LESILRRRF	1189	9	Human	Her2/neu	891	
GERLPOPPI	1190	9	Human	Her2/neu	938	
LEDDDMGDL	1191	9	Human	Her2/neu	1009	
EEYLVPQQG	1192	ģ	Human	Her2/neu	1021	
EEEAPRSPL	1193	9	Human	Her2/neu	1067	
EEAPRSPLA	1194	9	Human	Her2/neu	1068	
SEGAGSDVF	1195	9	Human	Her2/neu	1078	
PEYVNQPDV	1196	9	Human	Her2/neu	1137	
PEYLTPQGG	1197	9	Human	Her2/neu	1194	
PERGAPPST	1198	9	Human	Her2/neu	1228	
AENPEYLGL	1199	9	Human	Her2/neu	1243	
MELAALCRWG	1200	10	Human	Her2/neu	1	
LELTYLPTNA	1201	10	Human	Her2/neu	60	
QEVQGYVLIA		10	Human	Her2/neu	78	
FEDNYALAVL	1202	10	Human	Her2/neu	108	
TEILKGGVLI	1203	10	Human	Her2/neu	146	
	1204	10	Human	Her2/neu	206	
GESSEDCQSL SEDCQSLTRT	1205	10	Human	Her2/neu	209	
CELHCPALVT	1206	10	Human	Her2/neu	264	
	1207	10		Her2/neu	347	
MEHLREVRAV	1208		Human	Her2/neu	362	
QEFAGCKKIF	1209	10	Human	Her2/neu Her2/neu	400	
FETLEEITGY	1210	10	Human			
LEEITGYLYI	1211	10	Human	Her2/neu	403	
RELGSGLALI	1212	10	Human	Her2/neu	459	
PEDECVGEGL	1213	10	Human	Her2/neu	500	
QECVEECRVL	1214	10	Human	Her2/neu	538	
VEECRVLQGL	1215	10	Human	Her2/neu	541	
REYVNARHCL	1216	10	Human	Her2/neu	552	
PECQPQNGSV	1217	10	Human	Her2/neu	565	
EEGACQPCPI	1218	10	Human	Her2/neu	619	
QETELVEPLT	1219	10	Human	Her2/neu	692	
VEPLTPSGAM	1220	10	Human	Her2/neu	697	
KETELRKVKV	1221	10	Human	Her2/neu	716	
TELRKVKVLG	1222	10	Human	Her2/neu	718	
GENVKIPVAI	<u>1223</u>	10	Human	Her2/neu	743	

	one	н	ILA-B44 SUP	EKIYPE		
	SEQ ID					
Sequence	NO.	AA	Organism	Protein	Position	Analog
KEILDEAYVM	1224	10	Human	Her2/neu	765	
DEAYVMAGVG	1225	10	Human	Her2/neu	769	
DETEYHADGG	1226	10	Human	Her2/neu	873	
TEYHADGGKV	1227	10	Human	Her2/neu	875	
LESILRRRFT	1228	10	Human	Her2/neu	891	
REIPDLLEKG	1229	10	Human	Her2/neu	929	
SECRPRFREL	1230	10	Human	Her2/neu	963	
RELVSEFSRM	1231	10	Human	Her2/neu	970	
NEDLGPASPL	1232	10	Human	Her2/neu	991	
AEEYLVPQQG	1233	10	Human	Her2/neu	1020	
EEYLVPQQGF	1234	10	Human	Her2/neu	1021	
SEEEAPRSPL	1235	10	Human	Her2/neu	1066	
EEEAPRSPLA	1236	10	Human	Her2/neu	1067	
SETDGYVAPL	1237	10	Human	Her2/neu	1122	
PERGAPPSTF	1238	10	Human	Her2/neu	1228	
PEYLGLDVPV	1239	10	Human	Her2/neu	1246	
MELAALCRWGL	1240	11	Human	Her2/neu	1	
PETHLDMLRHL	1240	11	Human	Her2/neu	39	
RELOLRSLTEI	1241	11	Human	Her2/neu	138	
GESSEDCQSLT		11	Human	Her2/neu	206	
SEDCQSLTRTV	1243	11	Human	Her2/neu	209	
CELHCPALVTY	1244	11	Human	Her2/neu	264	
FESMPNPEGRY	1245	11	Human	Her2/neu	279	
	1246	11	Human	Her2/neu	331	
CEKCSKPCARV	1247	11	Human	Her2/neu	347	
MEHLREVRAVT	1248			Her2/neu	351	
REVRAVTSANI	1249	11 11	Human Human	Her2/neu	362	
QEFAGCKKIFG	1250			Her2/neu	400	
FETLEEITGYL	1251	11	Human			
EEITGYLYISA	1252	11	Human	Her2/neu	404	
GEGLACHQLCA	1253	11	Human	Her2/neu	506	
DEEGACQPCPI	<u>1254</u>	11	Human	Her2/neu	618	
AEQRASPLTSI	1255	11	Human	Her2/neu	644	
TELVEPLTPSG	<u>1256</u>	11	Human	Her2/neu	694	
KETELRKVKVL	1257	11	Human	Her2/neu	716	
KEILDEAYVMA	1258	11	Human	Her2/neu	765	
LEDVRLVHRDL	<u>1259</u>	11	Human	Her2/neu	836	
WELMTFGAKPY	<u>1260</u>	11	Human	Her2/neu	913	
GERLPQPPICT	<u>1261</u>	11	Human	Her2/neu	938	
SECRPRFRELV	<u>1262</u>	11	Human	Her2/neu	963	
RELVSEFSRMA	<u>1263</u>	11	Human	Her2/neu	970	
AEEYLVPQQGF	<u>1264</u>	11	Human	Her2/neu	1020	
EEYLVPQQGFF	<u>1265</u>	11	Human	Her2/neu	1021	
SEEEAPRSPLA	<u>1266</u>	11	Human	Her2/neu	1066	
SEGAGSDVFDG	<u>1267</u>	11	Human	Her2/neu	1078	
SETDGYVAPLT	1268	11	Human	Her2/neu	1122	
REGPLPAARPA	<u>1269</u>	11	Human	Her2/neu	1153	
VENPEYLTPQG	1270	11	Human	Her2/neu	1191	
PEYLTPQGGAA	1271	11	Human	Her2/neu	1194	
AENPEYLGLDV	1272	11	Human	Her2/neu	1243	
LELTYLPTNASL	1273	12	Human	Her2/neu	60	
RELQLRSLTEIL	1274	12	Human	Her2/neu	138	

		H	ILA-B44 SUP	ERTYPE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
PEGRYTFGASCV	1275	12	Human	Her2/neu	285	
LEEITGYLYISA	1276	12	Human	Her2/neu	403	
EEITGYLYISAW	1277	12	Human	Her2/neu	404	
PEADQCVACAH	12//	12	Human	Her2/neu	579	
Y	1278	12	Haman	Herzmea	517	
TELVEPLTPSGA	1279	12	Human	Her2/neu	694	
TEYHADGGKVPI	1280	12	Human	Her2/neu	875	
GERLPQPPICTI	1281	12	Human	Her2/neu	938	
AEEYLVPQQGFF	1282	12	Human	Her2/neu	1020	
PEGRYTFGASCV	1202	13	Human	Her2/neu	285	
T	1283					
CEKCSKPCARVC		13	Human	Her2/neu	331	
Y	1284					
MEHLREVRAVTS		13	Human	Her2/neu	347	
A	<u>1285</u>					
DECVGEGLACHQ		13	Human	Her2/neu	502	
L	<u>1286</u>		**	TT2/	565	
PECQPQNGSVTC F	1207	13	Human	Her2/neu	303	
RENTSPKANKEIL	1287 1288	13	Human	Her2/neu	756	
REIPDLLEKGERL	1289	13	Human	Her2/neu	929	
SEFSRMARDPQR	1203	13	Human	Her2/neu	974	
F	1290	13	Human	Herzmeu	214	
SEGAGSDVFDGD	1230	13	Human	Her2/neu	1078	
L	1291					
GEFGGYGSV	1292	9	Human	Histactranf	127	Α
LWQLNGRLEYTL		15	Human	IFN-B	21	Α
KDR	1293					
SEFQAAI	1294	7	Human	MAGE2	103	
SEYLQLV	<u>1295</u>	7	Human	MAGE2	155	
WEELSML	<u>1296</u>	7	Human	MAGE2	222	
GEPHISY	1297	7	Human	MAGE2	295	
LEARGEAL	1298	8	Human	MAGE2	16	
QEEEGPRM	1299	8	Human	MAGE2	90	
EEEGPRMF	1300	8	Human	MAGE2	91	
VELVHFLL	<u>1301</u>	8	Human	MAGE2	114	
AEMLESVL	1302	8	Human	MAGE2	133	
SEYLQLVF	1303	8	Human	MAGE2	155	
EEKIWEEL	1304	8	Human	MAGE2	218	
LEARGEALG	1305	9	Human	MAGE2	16	
GEALGLVGA	1306	9	Human	MAGE2	20	
QEEEGPRMF	1307	9	Human	MAGE2	90	
VELVHFLLL	<u>1308</u>	9	Human	MAGE2	114	
REPVTKAEM	1309	9	Human	MAGE2	127	
SEYLQLVFG	<u>1310</u>	9	Human	MAGE2	155	
PEEKIWEEL	1311	9	Human	MAGE2	217	
EELSMLEVF	1312	9	Human	MAGE2	223	
FEGREDSVF	1313	9	Human	MAGE2	231	
YEFLWGPRA	<u>1314</u>	9	Human	MAGE2	269	
EEGLEARGEA	1315	10	Human	MAGE2	13	
LEARGEALGL	<u>1316</u>	10	Human	MAGE2	16	
VEVTLGEVPA	1317	10	Human	MAGE2	46	
EEGPRMFPDL	<u>1318</u>	10	Human	MAGE2	92	

	CEC	h	ILA-B44 SUP	ERIYPE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
REPVTKAEML	1319	10	Human	MAGE2	127	
SEYLQLVFGI	1320	10	Human	MAGE2	155	
VEVVPISHLY	1321	10	Human	MAGE2	167	
EEKIWEELSM	1322	10	Human	MAGE2	218	
WEELSMLEVF	1323	10	Human	MAGE2	222	
FEGREDSVFA	1324	10	Human	MAGE2	231	
QENYLEYROV	1325	10	Human	MAGE2	252	
YEFLWGPRAL	1326	10	Human	MAGE2	269	
GEPHISYPPL	1327	10	Human	MAGE2	295	
EEGLEARGEAL	1328	11	Human	MAGE2	13	
LEARGEALGLV	1329	11	Human	MAGE2	16	
GEALGLVGAQA	1330	11	Human	MAGE2	20	
EEOOTASSSST	1331	11	Human	MAGE2	34	
VEVTLGEVPAA	1332	11	Human	MAGE2	46	
EEEGPRMFPDL	1333	11	Human	MAGE2	91	
SEFQAAISRKM	1334	11	Human	MAGE2	103	
VELVHFLLLKY	1335	11	Human	MAGE2	114	
LESVLRNCQDF	1336	11	Human	MAGE2	136	
VEVVPISHLYI	1337	11	Human	MAGE2	167	
IEGDCAPEEKI	1338	11	Human	MAGE2	211	
EEKIWEELSML	1339	11	Human	MAGE2	218	
EELSMLEVFEG	1340	11	Human	MAGE2	223	
LEVFEGREDSV	1341	11	Human	MAGE2	228	
YEFLWGPRALI	1342	11	Human	MAGE2	269	
EEQQTASSSSTL	1343	12	Human	MAGE2	34	
OEEEGPRMFPDL	1344	12	Human	MAGE2	90	
SEFQAAISRKMV	1345	12	Human	MAGE2	103	
LESVLRNCQDFF	1346	12	Human	MAGE2	136	
VEVVPISHLYIL	1347	12	Human	MAGE2	167	
EEGLEARGEALG L	1348	13	Human	MAGE2	13	
LEARGEALGLVG	15.10	13	Human	MAGE2	16	
A	1349					
LESEFQAAISRK		13	Human	MAGE2	101	
M	1350					
REPVTKAEMLES		13	Human	MAGE2	127	
V	<u>1351</u>			3.5.4.677.0		
SEYLQLVFGIEVV	1352	13	Human	MAGE2	155	
IEVVEVVPISHLY	1353	13	Human	MAGE2	164	
VEVVPISHLYILV	1354	13	Human	MAGE2	167	
MEVDPIGHLY	1355	10	Human	MAGE3	167	
EEEGPSTF	<u>1356</u>	8	Human	MAGE3	91	
AELVHFLL	<u>1357</u>	8	Human	MAGE3	114	
FEGREDSI	1358	8	Human	MAGE3	231	
QEAASSSST	1359	9	Human	MAGE3	36	
AELVHFLLL	1360	9	Human	MAGE3	114	
AEMLGSVVG	<u>1361</u>	9	Human	MAGE3	133	
EELSVLEVF	<u>1362</u>	9	Human	MAGE3	223	
FEGREDSIL	1363	9	Human	MAGE3	231	
QEAASSSSTL	<u>1364</u>	10	Human	MAGE3	36	
EEGPSTFPDL IELMEVDPIG	1365 1366	10 10	Human Human	MAGE3 MAGE3	92 164	

		H	ILA-B44 SUPI	ERTYPE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analo
MEVDPIGHLY	1367	10	Human	MAGE3	167	
EEKIWEELSV	1368	10	Human	MAGE3	218	
WEELSVLEVF	1369	10	Human	MAGE3	222	
FEGREDSILG	1370	10	Human	MAGE3	231	
EEEGPSTFPDL	1371	11	Human	MAGE3	91	
AELVHFLLLKY	1372	11	Human	MAGE3	114	
MEVDPIGHLYI	1373	11	Human	MAGE3	167	
REGDCAPEEKI	1374	11	Human	MAGE3	211	
EEKIWEELSVL		11	Human	MAGE3	218	
LEVFEGREDSI	1375	11	Human	MAGE3	228	
	1376	7			335	
RERFEMF	1377		Human	p53		
LEDSSGNL	<u>1378</u>	8	Human	p53	257	
GEYFTLQI	<u>1379</u>	8	Human	p53	325	
VEPPLSQET	1380	9	Human	p53	10	
PENNVLSPL	<u>1381</u>	9	Human	p53	27	
DEAPRMPEA	<u>1382</u>	9	Human	p53	61	
HERCSDSDG	1383	9	Human	p53	179	
VEGNLRVEY	1384	9	Human	p53	197	
VEYLDDRNT	1385	9	Human	p53	203	
LEDSSGNLL	1386	9	Human	p53	257	
RELNEALEL	1387	9	Human	p53	342	
NEALELKDA	1388	9	Human	p53	345	
LELKDAQAG	1389	9	Human	p53	348	
MEEPQSDPSV	1390	10	Human	p53	1	
VEPPLSQETF	1391	10	Human	p53	10	
QETFSDLWKL	1392	10	Human	p53	16	
IEQWFTEDPG	1393	10	Human	p53	50	
DEAPRMPEAA	1394	10	Human	p53	61	
HERCSDSDGL	1395	10	Human	p53	179	
VEGNLRVEYL	1396	10	Human	p53	197	
		10	Human	p53	203	
VEYLDDRNTF	1397				223	
PEVGSDCTTI	1398	10	Human	p53	257	
LEDSSGNLLG	1399	10	Human	p53		
FEVRVCACPG	1400	10	Human	p53	270	
TEEENLRKKG	<u>1401</u>	10	Human	p53	284	
GEPHHELPPG	<u>1402</u>	10	Human	p53	293	
GEYFTLQIRG	1403	10	Human	p53	325	
RERFEMFREL	1404	10	Human	p53	335	
FEMFRELNEA	1405	10	Human	p53	338	
QETFSDLWKLL	1406	11	Human	p53	16	
HERCSDSDGLA	1407	11	Human	p53	179	
YEPPEVGSDCT	1408	11	Human	p53	220	
HELPPGSTKRA	1409	11	Human	p53	297	
FEMFRELNEAL	1410	11	Human	p53	338	
NEALELKDAQA	1411	11	Human	p53	345	
TEDPGPDEAPRM	1412	12	Human	p53	55	
GEPHHELPPGST	1413	12	Human	p53	293	
DEAPRMPEAAPP		13	Human	p53	61	
V	1414	-		•		
YEPPEVGSDCTTI	1415	13	Human	p53	220	
		8	Human	unknown		

		H	LA-B44 SUP	ERTYPE		
	SEQ ID					
Sequence	NO.	AA	Organism	Protein	Position	Analog
SEIDLILGY	1417	9	Human	unknown		
AEIPTRVNY	1418	9	Human	unknown		
AEMGKFKFSY	1419	10	Human	unknown		
DEIGVIDLY	1420	9	Human	unknown		
AEMGKFKYSF	1421	10	Human	unknown		Α
SEAIHTFQY	1422	9	Human	unknown		
SEAIYTFQF	1423	9	Human	unknown		Α
AEGIVTGQY	1424	9	Human	unknown		
HETTYNSI	1425	8	Mouse	beta actin	275	Α
GELSYLNV	1426	8	Mouse	cathepsin D	255	
YEDTGKTI	1427	8	Mouse	p40 phox RNA	245	
YENDIEKKI	1428	9	Pf	CSP	375	

TABLE 22

		Н	LA-B44 SU	PERTYPE			
Sequence	SEQ ID NO.	B*1801	B*4001	B*4002	B*4402	B*4403	B*4501
SEAAYAKKI	983	8609	308	129	1685	61	287
GEFPYKAAA	984	286	170	3.9	746	2537	11
SEAPYKAIL	985	2258	29	8.8	440	170	262
SEAPKYAIL	986	2263	113	7.8	762	2260	479
AEFKYIAAV	987	48	2.8	6.5	28	21	4.9
AEIPYLAKY	988	116	7258	3159	44	30	668
AEIPKLAYF		1641	57	5.6	229	57	608
FPFDYAAAF	<u>989</u> 990	141	37	5.0	227	3,	000
FPFKYKAAF	990	155					
FPFKYAKAF		86					
FPFKYAAAF	992	16					
FAFKYAAAF	993	95					
FQFKYAAAF	994	22					
	995	187					
FDFKYAAAF	<u>996</u>	18281	271	23	183	164	1073
SENDRYRLL	<u>997</u>						
IEDPPYNSL	<u>998</u>	35457	10	688	15833	40075	18697
YEANGNLI	999	191	7.9	7.0	516	3085	10342
YEDLRVLSF	1000	20	67	71	24	212	18697
SDYEGRLI	1001	>24800	27150	86	851	228	10469
GEISPYPSL	<u>1002</u>	19361	24	1.8	3564	293	115
MDIDPYKEF	1003	169477	3700	382	21744	1949	2615
LDKGIKPY	1004	>100000	17884	468	>43192.49	19311	23609
ADLMGYIPL	1005	>7616.71	959	4.7	>21395.35	10292	>49000
LDPYARVAI	1006	>24409.45	>88888.89	372	>41628.96	>39766.08	>49000
AENLWVTVY	1007	155	1053	547	522	284	200
KENLWVTVY	1008	184	2738	373	308	306	6215
AEKLWVTVY	1009	286	18278	306	168	287	219
AENKWVTVY	1010	781	11303	534	294	540	297
AENLKVTVY	1011	138	7746	1075	253	487	9624
AENLWKTVY	1012	913	850	406	139	383	245
AENLWVKVY	1013	2735	1482	1696	708	105	132
AENLWVTKY	1014	511	1010	1998	355	1064	201
AENLWVTVK	1015	29464	853	2004	6305	2133	186
FENLWVTVY	1016	59	943	1336	4179	1312	21403
VENLWVTVY	1017	25	5499	5586	13454	4856	15654
PENLWVTVY	1018	190	>72727.27	>154545.45	>167272.73	>425000	>49000
NENLWVTVY	1019	38	>72727.27	11774	453	224	1668
DENLWYTVY	1020	26	>72727.27	41098	4589	988	49000
TENLWVTVY	1021	14	14040	1415	291	364	5296
YENLWVTVY	1022	29	552	324	640	369	10701
ATNLWVTVY	1023	17615	487	>154545.45	8912	>43037.97	>49000
AEFLWVTVY	1024	131	183	240	1013	156	472
AEVLWVTVY	1025	142	1549	436	1520	390	1244
AEPLWVTVY	1026	310	1727	2484	1322	96	1384
AEDLWVTVY	1027	354	423	3521	2329	469	1845
AENLWVTVY	1028	122	1581	552	308	132	301
AETLWVTVY	1029	199	1052	198	501	221	774
AENFWVTVY	1030	182	1394	542	171	268	289
AENVWVTVY	1031	262	2238	386	1112	744	737
AENPWVTVY	1031	27	843	224	18	53	202

		н	A-B44 SU	PERTYPE			
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AENDWVTVY	1033	324	954	742	96	165	365
AENNWVTVY	1034	167	1161	357	214	162	99
AENTWVTVY	1035	213	1451	1793	386	166	442
AENLFVTVY	1036	29	970	334	357	125	232
AENLVVTVY	1037	62	876	1344	1030	203	718
AENLPVTVY	1038	20	205	566	356	126	246
AENLDVTVY	1039	517	220	12081	673	340	1291
AENLNVTVY	1040	198	564	3544	447	358	2445
AENLTVTVY	1041	153	689	1269	327	208	793
AENLWFTVY	1042	360	699	668	227	62	90
AENLWLTVY	1043	666	1702	884	647	226	227
AENLWPTVY	1044	661	690	688	157	50	116
AENLWDTVY	1045	775	1145	2090	414	68	263
AENLWNTVY	1046	336	1338	957	66	81	257
AENLWTTVY	1047	196	246	625	51	50	118
AENLWVFVY	1048	242	857	375	348	310	237
AENLWVVVY	1049	326	2728	1688	599	632	468
AENLWVPVY	1050	303	175	183	96	47	106
AENLWVDVY	1051	415	700	3440	334	92	242
AENLWVNVY	1052	317	1156	952	159	76	266
AENLWVSVY	1053	232	1251	1347	351	178	292
AENLWVTFY	1054	1299	1201	295	124	222	347
AENLWVTLY	1055	392	463	731	199	119	349
AENLWVTPY	1056	41	274	189	127	44	122
AENLWYTDY	1057	1001	930	1208	191	103	328
AENLWVTNY	1058	730	865	948	149	74	215
AENLWYTTY	1059	28	280	191	37	26	48
AENLWVTVA	1060	9689	557	4.8	1543	296	9.1
AENLWVTVC	1061	178026	157	1425	5593	2267	146
AENLWYTVE	1062	>258333.33	3888	1362	8910	2573	246
AENLWVTVF	1063	365	162	20	346	162	262
AENLWVTVG	1064	39743	861	47	1812	245	35
AENLWVTVH	1065	16516	493	151	966	387	120
AENLWVTVI	1066	11224	14	7.3	237	88	54
AENLWVTVL	1067	6198	14	13	68	208	114
AENLWVTVM	1068	508	13	6.1	195	35	50
AENLWVTVN	1069	129167	6701	481	2623	414	169
AENLWVTVP	1070	38441	9711	339	7715	2473	187
AENLWVTVQ	1070	49640	522	85	1223	188	100
AENLWVTVR	1071	32979	1246	1744	4857	1474	233
AENLWVTVS	1072	25726	2163	103	4221	417	34
AENLWVTVT	1073	12331	947	7.8	2696	343	10
AENLWVTVV	1074	10709	84	19	5757	1432	35
AENLWVTVW		22610	1304	135	423	324	204
AENLWVTVY	1076	51	1358	90	66	43	68
AENLYVTVF	1077	61	17	3.1	39	47	69
TEPAAVGVGAV	1078	>8115.18	930	391	1938	459	8235
AEPAAEGV	1079	>8115.18 >8115.18	2070	2675	>22604.42	402	6590
AEPAAEGV AEPAAEGVGA	1080	>8115.18 >8115.18	4116	1655	>22604.42	402 >11447.81	104
AEPAAEGVGA AEPAAEGVGAV	1081	>8115.18 >8611.11	20364	242	>22604.42	>11447.81	1499
QEEEEVGFPV	1082	>8611.11	13117	2596	15203	>11447.81	86
EEEEVGFPV	1083 1084	3691	3340	417	7440	10313	37

		Н	LA-B44 SU	PERTYPE			
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EEEVGFPV	1085	427	9578	2605	6372	>10461.54	227
EEVGFPVRPQV	1086	>22794.12	9905	108	23777	6553	808
DEEVGFPV	1087	7.1	>32000	4260	9305	>10461.54	916
KEKGGLDGL	1088	>22794.12	55	174	>81415.93	>10461.54	9926
KEKGGLDGLI	1089	>22794.12	843	233	14726	3626	9986
QEILDLWV	1090	>22794.12	142	1717	>81415.93	5919	5504
QEILDLWVY	1091	52	740	4522	264	172	6261
AETFYVDGA	1092	>6709.96	21630	1923	>21198.16	6924	38
EEKPRTLHDL	1093	>81578.95	36208	34027	15236	30010	419
NEILIRCII	1094	5672	291	59	2722	258	3248
QEKKRHVDL	1095	7.3	15984	63093	443	211	12613
AEGKEVLL	1096	11455	1311	5303	17268	129	14165
QELFIPNI	1097	127	5815	147	752	8.5	1319
QELFISNI	1098	889	6396	1175	2282	70	1172
TEKNSGLY	1099	211	9851	7117	1868	605	10248
AELPKPSI	1100	7423	6697	131	1164	19	2608
PEAQNTTY	1101	149	2594	2437	2204	76	3255
IESTPFNVA	1102	69	1234	66	18749	0.97	15
AEGKEVLLL	1103	1080	72	147	178	1.7	199
EEATGQFRV	1104	805	5563	470	1691	95	18
VEDKDAVAF	1105	94	121	1583	1661	1443	21204
CEPETQDAT	1106	4009	3646	410	23421	50	97
PETQDATYL	1107	9473	1240	33745	>34586.47	301	13430
CETQNPVSA	1108	73	7016	261	20023	10.0	15
QELFIPNIT	1109	125	4361	172	1217	3.0	18
AEPPKPFIT	1110	12850	7067	7170	>34586.47	232	1813
VEDEDAVAL	1111	840	I 1	2665	30667	51	27810
CEPEIQNTT	1112	6889	5709	3081	31834	120	2732
PEIQNTTYL	1113	923	138	2786	16816	231	1825
YECGIQNEL	1114	82	71	53	452	5.3	855
QELFISNIT	1115	530	6571	58	2334	3.9	80
TEKNSGLYT	1116	1113	7522	3195	10097	101	1963
AEGKEVLLLV	1117	5135	1019	408	479	8.6	994
KEVLLLVHNL	1118	893	3.1	4.4	414	2.3	2512
GERVDGNRQI	1119	9395	1933	369	3900	13	19464
REIIYPNASL	1120	74 I	2.3	7.5	374	1.7	954
NEEATGQFRV	1121	998	29086	22678	4365	471	405
EEATGQFRVY	1122	64	>33333.33	55956	29	1041	1374
GENLNLSCHA	1123	14373	1341	357	8610	5.3	271
QELFIPNITV	1124	81	121	27	93	2.6	14
CEPEIQNTTY	1125	1459	>10322.58	35697	49	14596	43739
PEIQNTTYLW	1126	819	3301	9423	13	6173	10011
CEPEAQNTTY	1127	9525	>12903.23	>48571.43	61	>4268.68	17330
PEAQNTTYLW	1128	17082	>9248.55	>12592.59	27	21243	>28654.97
MESPSAPPHRW	1129	12	943	1915	5.3	41	359
IESTPFNVAEG	1130	87	1074	352	89	8.7	84
GERVDGNRQII	1131	764	278	18	871	1.3	27084
REIIYPNASLL	1132	1788	2.4	12	57	0.38	1777
NEEATGQFRVY	1133	7.7	3252	999	9.6	69	3986
CEPETQDATYL	1134	831	311	3388	398	807	62150
GENLNLSCHAA	1135	7838	4557	63	1907	9.0	32
CEPEIQNTTYL	1136	129	287	1603	1245	60	11981

		Ė	ILA-B44 SUF	PERTYPE			
	SEQ ID	B*1801	B*4001	B*4002	B*4402	B*4403	B*4501
Sequence PEIONTTYLWW	NO.	172	749	1045	17	227	1365
YECGIONELSV	1137	9.2	33	26	1714	0.46	155
NELSVDHSDPV	1138	49	2554	1128	1615	38	78
	1139	962	2184	11723	3419	131	2450
CEPEAQNTTYL	1140	147	2096	3090	121	79	2005
PEAQNTTYLWW	1141	644	1808	1539	481	93	994
PEIQNTTYLWWV	1142	20	1694	646	5.1	73	3.3
PEAQNTTYLWWV	1143	84			7.9	409	1243
CEPEIQNTTYLW W	1144	84	858	3168	7.9	409	1243
AEMGKGSFKY	1145	1618	6427	3820	112	90	305
SEDCOSL	1146	18245	2691	14258	8248	431	19225
REVRAVT	1147	8564	3136	725	31615	29	23544
FETLEE1	1148	1518	7621	2110	42991	69	67957
TELVEPL	1149	162	14164	1258	8854	66	>148484.85
SECRPRF	1150	926	18181	1157	852	48	8856
PETHLDML	1151	1954	8387	6118	>17523.81	83	20257
QEVQGYVL	1152	3.4	28	5.0	1210	0.92	33
RELQLRSL	1153	42	49	5.9	2025	0.62	1372
CELHCPAL	1154	150	871	259	4361	39	30089
LEEITGYL	1155	242	830	1805	5913	403	35502
EEITGYLY	1156	20	5713	1223	11	83	238
DECVGEGL	1157	49	4864	481	938	34	14244
AEORASPL	1157	16	73	13	211	0.38	120
KEILDEAY	1159	82	921	430	1081	74	2646
EEAPRSPL	1160	1191	3489	1611	1593	171	1926
SEDPTVPL	1161	103	71	161	12267	2.0	308
MELAALCRW		7.0	4833	138	16	9.9	1183
QEVQGYVLI	1162	77	206	39	30	0.50	96
FEDNYALAV	1163	12	34	5.1	13470	0.17	131
RELQLRSLT	1164	638	316	13	465	0.20	162
TEILKGGVL	1165	125	30	14	1377	0.28	2480
HEOCAAGCT	1166	1995	42164	7377	19048	178	2974
CELHCPALV	1167	136	4805	319	2308	52	1110
FESMPNPEG	1168	6068	30237	59	16458	14	155
QEVTAEDGT	1169	5207	31081	3122	7886	66	1843
CEKCSKPCA	1170	3740	27386	2703	19957	342	8007
MEHLREVRA	1171	233	44754	386	38	3.2	19
REVRAVTSA	1172	626	44734	0.71	3160	0.18	9.3
QEFAGCKKI	1173	1120	736	131	81	44	2684
EEITGYLYI	1174	86	906	916	12	121	94
RELGSGLAL	1175	359	3.7	0.85	457	0.97	2262
	<u>1176</u>	13766	187	88	112	11	340
GEGLACHQL	1177	15799	8755	1664	7150	210	4542
QECVEECRV	1178	1528	8947	7622	14202	305	20142
VEECRVLQG	1179	1528 890	8947 7076	2029	717	434	1185
EECRVLQGL	1180		874		103	1.8	1185
AEQRASPLT	1181	346		183			
QETELVEPL	1182	12	62	85	681	3.5	1232
VEPLTPSGA	1183	7321	>9638.55	11	8516	191	17037
TELRKVKVL	1184	1514	4698	54	2128	2.5	14147
GENVKIPVA	<u>1185</u>	10755	14510	7.5	20309	2.7	7.0
KEILDEAYV	<u>1186</u>	1358	62	146	6466	8.4	42
DEAYVMAGV	<u>1187</u>	58	5327	1245	8006	138	161

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Sequence	ID NO.	B*1801	B*4001	B*4002	B*4402	B*4403	B*4501		
DETEYHADG	1188	159	>11940.3	>65384.62	>24403.18	1397	13353		
LESILRRRF	1189	29	>11940.3	3475	4.7	101	12918		
GERLPOPPI	1190	62	71	15	63	1.1	15		
LEDDDMGDL	1191	191	556	351	947	900	6251		
EEYLVPOOG	1192	66	10344	136	651	126	131		
EEEAPRSPL	1193	902	4490	2881	342	362	307		
EEAPRSPLA	1194	486	10707	4900	180	294	4.5		
SEGAGSDVF	1195	74	5627	6525	69	192	6960		
PEYVNOPDV	1196	831	3437	1581	1109	48	2536		
PEYLTPOGG	1197	1456	18951	13860	6532	284	18990		
PERGAPPST	1198	385	4744	7679	1116	178	7767		
AENPEYLGL	1199	17	81	271	44	2.5	155		
MELAALCRWG	1200	102	8684	1840	5.7	135	408		
LELTYLPTNA	1201	332	325	10.4	6428	3.1	24		
QEVQGYVLIA	1201	61	772	64	1871	15	11		
FEDNYALAVL	1202	321	6.2	48	2844	3.8	3095		
TEILKGGVLI	1204	1021	241	294	24	21	7600		
GESSEDCOSL	1204	138636	8.1	23	427	5.1	2491		
SEDCQSLTRT	1205	335	8550	11529	518	2857	4726		
CELHCPALVT	1200	80	>9248.55	65	933	18	477		
MEHLREVRAV	1207	72	20684	160	180	13	140		
OEFAGCKKIF	1208	53	3686	12	4.0	3.6	115		
FETLEEITGY	1210	671	53363	36302	262	1679	>28488.37		
LEEITGYLYI		143	914	2996	222	143	1488		
RELGSGLALI	1211 1212	4810	22	4.4	32	0.78	173		
PEDECVGEGL	1212	1257	278	257	6331	49	24019		
QECVEECRVL	1213	315	444	399	606	22	2863		
VEECRVLOGL	1214	270	227	5815	237	189	16094		
REYVNARHCL	1215	1327	39	4.8	106	0.97	126		
PECQPQNGSV	1216	7962	35957	20374	12964	472	>28488.37		
EEGACOPCP1		119	40113	340	52	80	401		
OETELVEPLT	1218 1219	15	293	338	1619	13	288		
VEPLTPSGAM		4649	1667	584	4368	108	20167		
KETELRKVKV	1220	11925	26700	68	2936	4.5	1603		
TELRKVKVLG	1221	721	20700	601	3650	14	12816		
GENVKIPVAI	1222	563	314	28	230	6.7	198		
KEILDEAYVM	1223	0.14	10	153	35	7.5	234		
DEAYVMAGVG	1224	122	203	154	4033	4102	218		
DETEYHADGG	1225	613	45291	16801	3891	269	29025		
TEYHADGGKV	1226	239	5246	2003	2911	15	1571		
LESILRRRFT	1227	82	28476	1189	34	87	2251		
REIPDLLEKG	1228	649	4493	814	1270	13	1977		
SECRPRFREL	1229	80	307	18	11	0.20	25		
RELVSEFSRM	1230	9.1	28	4.3	33	0.12	1726		
NEDLGPASPL	1231	107	281	150	40	6.0	231		
AEEYLVPQQG	1232	723	66699	24424	417	479	127		
EEYLVPQQG	1233	2.1	26569	2551	6.9	11	73		
SEEEAPRSPL	1234	151	155	2331	37	8.4	73 84		
EEEAPRSPLA	1235	6611	49549	38943	425	960	14		
SETDGYVAPL	1236	94	214	184	386	2.4	302		
PERGAPPSTF	1237	1062	14884	3437	6871	2.4	15700		
PEYLGLDVPV	1238	613	352	3437	1371	1.7	610		
LE I LULDALA	<u>1239</u>	013	334	33	1371	1.7	010		

		I	ILA-B44 SUF	ERTYPE			
	SEQ ID	D41001	B*4001	B*4002	B*4402	B*4403	B*4501
Sequence MELAALCRWGL	NO.	B*1801 6.4	24	30	17	0.92	116
PETHLDMLRHL	1240	1322	700	2971	11534	70	4329
	1241	261	2.8	3.7	125	0.99	269
RELQLRSLTEI	1242	742	48	180	14386	40	2158
GESSEDCQSLT	1243		48	311	943	21	10
SEDCQSLTRTV	1244	101				21 89	2779
CELHCPALVTY	<u>1245</u>	12	3469	3198	140		
FESMPNPEGRY	<u>1246</u>	74	3666	3533	59	70	1394
CEKCSKPCARV	<u>1247</u>	1167	4103	2079	9594	101	1561
MEHLREVRAVT	<u>1248</u>	1064	3614	2207	795	111	74
REVRAVTSANI	<u>1249</u>	4491	17	30	1680	1.8	421
QEFAGCKKIFG	1250	211	314	477	37	2.1	138
FETLEEITGYL	1251	133	78	649	7490	42	2200
EEITGYLYISA	1252	0.94	1440	52	4.5	2.1	0.9
GEGLACHQLCA	1253	62	39	97	159	2.7	196
DEEGACQPCPI	1254	451	5517	7293	968	438	1323
AEQRASPLTSI	1255	467	19	58	5.1	2.5	11
TELVEPLTPSG	1256	601	2978	3703	>21052.63	269	14079
KETELRKVKVL	1257	9529	2973	1868	7136	71	12237
KEILDEAYVMA	1258	731	252	95	11514	64	123
LEDVRLVHRDL	1259	729	325	641	818	59	2382
WELMTFGAKPY	1260	13	509	778	24	75	1216
GERLPOPPICT	1261	12486	24270	23	9094	3.9	15
SECRPRFRELV	1262	1996	3673	121	927	18	118
RELVSEFSRMA	1263	168	389	143	2613	3.5	32
AEEYLVPOOGF	1264	125	584	1831	21	99	268
EEYLVPQQGFF		94	4291	1695	78	168	154
SEEEAPRSPLA	1265	1318	3604	5110	8550	158	27
	1266				374	286	3008
SEGAGSDVFDG	<u>1267</u>	928	3751	5695			
SETDGYVAPLT	<u>1268</u>	66	125	224	1225	2.2	45
REGPLPAARPA	<u>1269</u>	157	543	78	32906	4.2	347
VENPEYLTPQG	<u>1270</u>	8386	56393	42593	17337	11	4188
PEYLTPQGGAA	<u>1271</u>	1724	41026	200	>17829.46	354	1382
AENPEYLGLDV	<u>1272</u>	11934	28	139	69	3.0	24
LELTYLPTNASL	1273	12	25	102	386	6.8	11
RELQLRSLTEIL	1274	5954	151	600	3778	1.1	1371
PEGRYTFGASCV	1275	4071	2.9	4.4	778		116
LEEITGYLYISA	1276	209	28	31	263	18	694
EEITGYLYISAW	1277	746	478	1800	252		1492
PEADQCVACAHY	1278	901	4050	5127	213		463
TELVEPLTPSGA	1279	236	2059	59	2132		206
TEYHADGGKVPI	1280	680	22	4.4	2177		61
GERLPQPPICTI	1281	17769	162	3.9	292		2.5
AEEYLVPQQGFF	1282	144	228	45	16		13
PEGRYTFGASCVT	1283	5228	3793	737	1419	267	673
CEKCSKPCARVC Y	1284	701	>53333.33	406	302	44	1315
MEHLREVRAVTS A	1285	70	669	72	144	18	12
DECVGEGLACHQ L	1286	464	2635	3668	2544	212	2063
PECQPQNGSVTCF	1287	6293	381	5338	3564	375	>22374.
RENTSPKANKEIL	1288	7750	3.7	77	>2540.03	3.9	1510
REIPDLLEKGERL	1289	7636	40	136	3050	16	2710

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	SEQ ID									
Sequence	NO.	B*1801	B*4001	B*4002	B*4402	B*4403	B*4501			
SEFSRMARDPQRF	1290	61	350	57	23	12	247			
SEGAGSDVFDGD		5172	45	2059	1303	711	2458			
L	1291				2226		40			
GEFGGYGSV	<u>1292</u>	307	112	6.4	2335	534	40			
LWQLNGRLEYTL KDR	1293					0.11				
SEFQAAI	1294	181	6830	779	2660	33	9597			
SEYLQLV	1295	1375	7777	658	733	21	930			
WEELSML	1296	1288	781	740	>28482.97	151	82009			
GEPHISY	1297	8833	12272	6716	36116	272	>33333.33			
LEARGEAL	1298	163	99	65	29495	2.9	31463			
OEEEGPRM	1299	298	11598	1608	19255	118	6730			
EEEGPRMF	1300	723	12281	32093	2406	213	943			
VELVHFLL	1301	5.0	69	31	3322	1.2	2427			
AEMLESVL	1302	968	14	31	327	0.88	302			
SEYLQLVF	1302	0.97	765	6.0	284	0.70	122			
EEKIWEEL	1303	753	9084	2599	98976	104	171			
LEARGEALG		155	1161	3006	11018	24	2688			
GEALGLVGA	1305	9529	2832	34	6134	2.2	17			
OEEEGPRMF	1306	414	918	7747	237	409	2171			
VELVHFLLL	1307	71	79	31	579	3.1	1129			
REPVTKAEM	1308	60	373	284	896	4.5	832			
SEYLQLVFG	1309	18	8890	421	271	19	113			
PEEKIWEEL	1310	577	19449	3908	1029	235	17345			
EELSMLEVF	1311	1.4	16436	252	22	2.8	1013			
FEGREDSVF	1312	9.8	2366	348	221	13	3339			
YEFLWGPRA	1313	5.3	249	5.2	2355	1.1	241			
EEGLEARGEA	1314	3.3 1077	3434	3227	2333	302	30			
	1315	81	184	277	2275	4.1	964			
LEARGEALGL	<u>1316</u>	14	371	31	3801	0.52	15			
VEVTLGEVPA	1317					13	42			
EEGPRMFPDL	1318	128	4438	486	95	41	917			
REPVTKAEML	<u>1319</u>	88	23	264	84	0.84				
SEYLQLVFGI	<u>1320</u>	2.2	20	6.1	3.7		4.4			
VEVVPISHLY	<u>1321</u>	20	11522	4385	13	1225	4885			
EEKIWEELSM	1322	17	21450	477	46	19	107			
WEELSMLEVF	1323	0.14	463	30	15	15	290			
FEGREDSVFA	1324	178	>10062.89	4775	6879	192	503			
QENYLEYRQV	1325	118	493	102	17	16	27			
YEFLWGPRAL	1326		8.5	0.97	130	0.72	753			
GEPHISYPPL	1327	2612	7.0	2.9	1200	0.71	380			
EEGLEARGEAL	1328	179	300	578	2630	19	1812			
LEARGEALGLV	1329	158	198	345	>17829.46	13	1912			
GEALGLVGAQA	1330	877	4293	52	3575	1.4	28			
EEQQTASSSST	<u>1331</u>	752	4040	41162	5910	1552	134			
VEVTLGEVPAA	1332	124	25216	919	>23469.39	44	1583			
EEEGPRMFPDL	1333	1011	2646	3470	3273	131	209			
SEFQAAISRKM	1334	7.0	345	107	88	1.2	161			
VELVHFLLLKY	1335	52	550	294	1551	49	1790			
LESVLRNCQDF	1336	64	5409	3458	209	76	15241			
VEVVPISHLYI	1337	97	135	146	335	7.2	3788			
IEGDCAPEEKI	1338	844	27827	32058	2627	486	183			
EEKIWEELSML	<u>1339</u>	1641	4978	20625	1862	375	181			

HLA-B44 SUPERTYPE									
	SEQ								
Sequence	ID NO.	B*1801	B*4001	B*4002	B*4402	B*4403	B*4501		
EELSMLEVFEG	1340	1.5	24061	294	4.6	23	163		
LEVFEGREDSV	1340	639	2624	367	>21296.3	46	29449		
YEFLWGPRALI	1341	5.2	4.1	2.8	92	0.59	450		
EEQQTASSSSTL	1342	7259	166	526	57	981	15		
QEEEGPRMFPDL	1344	3595	394	1330	1643	,,,,	120		
SEFQAAISRKMV	1344	43	161	29	25		21		
LESVLRNCODFF	1345	56	55	356	184	24	1993		
VEVVPISHLYIL	1346	266	3.4	16	486	4.0	1182		
EEGLEARGEALG	1347	10416	1769	5143	196	118	1673		
L	1348	10410	1705	3143	150	110	1075		
LEARGEALGLVG		347	20	48	2575	2.2	116		
Α	1349		***		242				
LESEFQAAISRKM	<u>1350</u>	49	310	72	242	14	22		
REPVTKAEMLES V	1351	5531	337	411	4546	21	1507		
SEYLOLVFGIEVV	1352	9.7	23	4.5	144	5.4	6.6		
IEVVEVVPISHLY	1353	79	162	245	52	125	106		
VEVVPISHLYILV	1354	92	93	47	270	51	112		
MEVDPIGHLY	1355	13	209	334	13	28	228		
EEEGPSTF	1356	216	1008	435	3933	27	1819		
AELVHFLL		120	71	6.8	1074	0.16	452		
FEGREDS1	1357	927	718	127	7708	13	2291		
QEAASSSST	1358	1422	23469	1480	9593	41	110		
AELVHFLLL	1359 1360	160	25	3.1	33	0.94	141		
AEMLGSVVG		96	1899	109	27	1.6	11		
EELSVLEVF	1361 1362	7.3	10215	3314	61	12	2120		
FEGREDSIL	1363	1091	51	439	1925	11	>27071.82		
QEAASSSSTL	1364	171	49	47	56	13	287		
EEGPSTFPDL	1365	158	655	591	198	127	128		
IELMEVDPIG	1366	194	6592	5325	222	>16306.95	7604		
MEVDPIGHLY	1367	15	617	625	11	99	169		
EEKIWEELSV	1368	73	8947	79	396	17	17		
WEELSVLEVF	1369	1.7	75	37	14	13	1701		
FEGREDSILG	1370	229	940	4361	8534	172	20261		
EEEGPSTFPDL	1370	935	431	2120	2685	102	158		
AELVHFLLLKY	1371	153	32	39	178	1.6	670		
MEVDPIGHLYI	1372	9.8	34	16	64	0.91	95		
REGDCAPEEKI	1374	973	2418	830	4038	42	146		
EEKIWEELSVL	1374	133	152	1255	1416	58	218		
LEVFEGREDSI	1376	4745	206	512	20963	69	>31012.66		
RERFEMF	1377	180	4079	1907	25488	108	20048		
LEDSSGNL		17736	782	362	42791	211	15946		
GEYFTLOI	1378 1379	7774	112	60	3511	1.0	261		
VEPPLSQET	1380	8302	17052	20808	3186	236	29270		
PENNVLSPL	1381	1150	1261	718	11174	8.8	>27071.82		
DEAPRMPEA	1382	84	9092	4577	6448	98	10.0		
HERCSDSDG	1382	1118	2367	38636	19328	208	13390		
VEGNLRVEY	1383 1384	832	12752	67730	142	2583	39059		
VEYLDDRNT	1384	1442	36833	35854	10071	157	13503		
LEDSSGNLL		1140	43	2771	4656	43	26134		
RELNEALEL	1386	3000	43 15	30	525	1.1	3337		
NEALELKDA	1387	1925	3887	27585	4270	1582	129		
LELKDAQAG	1388 1389	451	18706	3659	17293	30	1989		
LLLINDAQAU	1389	731	10700	3033	17273	50	1707		

		Н	LA-B44 SUI	PERTYPE			
	SEQ						
Sequence	ID NO.	B*1801	B*4001	B*4002	B*4402	B*4403	B*4501
MEEPQSDPSV	1390	12157	3802	16536	1927	816	175
VEPPLSQETF	1391	814	>37209.3	21732	406	525	>24019.61
QETFSDLWKL	1392	736	199	255	39	14	901
IEQWFTEDPG	1393	151	1250	2114	5595	142	197
DEAPRMPEAA	1394	121	3941	8444	2594	1037	100
HERCSDSDGL	1395	139	171	61	1468	6.0	1723
VEGNLRVEYL	1396	104	481	2565	1963	22	15189
VEYLDDRNTF	1397	0.94	501	37	32	1.4	3601
PEVGSDCTTI	1398	611	4552	248	2293	2046	22487
LEDSSGNLLG	1399	103	531	697	7905	153	19256
FEVRVCACPG	1400	64	2043	4.9	180	0.76	1872
TEEENLRKKG	1401	74966	>37209.3	11858	>23589.74	315	30635
GEPHHELPPG	1402	108	3323	1888	11728	4.4	20
GEYFTLQIRG	1403	108	88	19	2452	3.9	157
RERFEMFREL	1404	83	29	17	17	0.34	422
FEMFRELNEA	1405	127	3207	223	952	2.0	208
QETFSDLWKLL	1406	4158	3366	740	631	168	1218
HERCSDSDGLA	1407	1408	4879	1915	>20956.72	96	186
YEPPEVGSDCT	1408	16872	4529	125	13349	12712	16034
HELPPGSTKRA	1409	6034	3974	3255	47077	189	1472
FEMFRELNEAL	1410	475	17	8.8	748	1.1	1352
NEALELKDAQA	1411	742	6235	5071	>20956.72	949	53
TEDPGPDEAPRM	1412	888	327	893	2053	161	1676
GEPHHELPPGST	1413	6822	24342	4631	6581	252	169
DEAPRMPEAAPP		427	>48484.85	7258	>2762.76	1376	19
V	1414						
YEPPEVGSDCTTI	1415	8796	2699	1540	>2740.54	253	>20000
RERRDNYV	<u>1416</u>	>73809.52	71554	62	>67647.06	>34517.77	34648
SEIDLILGY	1417	3.0	285	140	4.8	8.5	397
AEIPTRVNY	1418	1691	7826	5443	333	23	1286
AEMGKFKFSY	1419	1517	2941	622	146	28	283
DEIGVIDLY	1420	11	>114285.71		707	212	>49000
AEMGKFKYSF	1421	155	113	3.8	18	31	186
SEAIHTFQY	1422	25	2895	1802	18	16	1078
SEAIYTFQF	1423	5.7	967	39	4.8	20	293
AEGIVTGQY	1424	7176	6462	1528	255	12	418
HETTYNSI	1425	1644	251	336	616	23959	6608
GELSYLNV	1426	>24800	4856	100	19013	23735	784
YEDTGKTI	1427	13997	794	83	7911	2177	49000
YENDIEKKI	1428	30992	1156	145	1725	371	

TABLE 23

		ILA-DQ	SUPERTYPES			
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
AAAKAAAAAAYAA	1429	13	Artificial sequenc	e		Α
(44)YAAAAAKAAA	1430	13	Artificial sequenc	e		Α
AAFAAAKTAAAFA	1431	13	Artificial sequenc	e		Α
YAAFAAAKTAAAFA	1432	14	Artificial sequenc	e		Α
YAAFAAAKTAAAFA	1433	14	Artificial sequenc	e		
АНААНААНААНА		16	HA			Α
A	<u>1434</u>					
VLERYLLEAKEAENI	1435	15	Human	EPO	11	
VPDTKVNFYAWKRME	1436	15	Human	EPO	41	
WKRMEVGQQAVEVWQ	1437	15	Human	EPO	51	
VGQQAVEVWQGLALL	1438	15	Human	EPO	56	
VEVWQGLALLSEAVL	1439	15	Human	EPO	61	
GLALLSEAVLRGQAL	1440	15	Human	EPO	66	
SEAVLRGQALLVNSS	1441	15	Human	EPO	71	
RGQALLVNSSQPWEP	1442	15	Human	EPO	76	
LQLHVDKAVSGLRSL	1443	15	Human	EPO	91	
KEAISPPDAASAAPL	1444	15	Human	EPO	116	
PPDAASAAPLRTITA	1445	15	Human	EPO	121	
SAAPLRTITADTFRK	1446	15	Human	EPO	126	
EAENITTGTAEHTSL	1447	15	Human	EPO	21	Α
RLFDNASLRAHRLHQ		15	Human	Growth	8	
	1448			hormone		
QLAFDTYQEFEEAYI		15	Human	Growth	22	
IGI I I IOGUI ERVOE	<u>1449</u>	15	Human	hormone Growth	78	
ISLLLIQSWLEPVQF	1450	13	riuman	hormone	70	
NSLVYGASDSNVYDL	1.50	15	Human	Growth	99	
	1451			hormone		
SDSNVYDLLKDLEEG		15	Human	Growth	106	
KIROOL ARI BEGEDODDA	1452	10	TT	hormone	260	
KIFGSLAFLPESFDGDPA	<u>1453</u>	18	Human	Her2/neu	369	
CLKDRRNFDIPEEIK	<u>1454</u>	15	Human	IFN-B	31	
QLQQFQKEDAAVTIY	<u>1455</u>	15	Human	IFN-B	46	
QKEDAAVTIYEMLQN	<u>1456</u>	15	Human	IFN-B	51	
STGWNETIVENLLAN	1457	15	Human	IFN-B	76	
ETIVENLLANVYHQR	1458	15	Human	IFN-B	81	
KEDSHCAWTIVRVEI	1459	15	Human	IFN-B	136	
MSYNLLGFLQRSSNT	1460	15	Human	IFN-B	1	Α
QHLCGSHLVEALYLV		15	Human	Insulin	4	
	1461			beta chain		
GSHLVEALYLVCGER	<u>1461</u>	15	Human	Insulin	8	
GOILVERLIEVCOER		15		beta		
	1462			chain		
GSDLVEALYLVCGER		15	Human	Insulin	8	Α
	1462			beta		
VEALYLVCGERGFLY	1463	15	Human	chain Insulin	12	Α
VEAL I EV COEROI LI	<u>1464</u>	13	raman	mounn	12	^

	HLA-DQ SUPERTYPES										
	SEQ ID										
Sequence	NO.	AA	Organism	Protein	Position	Analog					
				beta							
				chain							
VEALYLVTGERGFFY		15	Human	Insulin beta	12	Α					
	1465			chain							
IDVWLGGLAENFLPY	1100	15	Human	thyroid	632						
	1466			perox							
IDVWLGGLAYNFLPY		15	Human	thyroid	632	Α					
IDVIVI CCI ALNEI DV	<u>1467</u>	15	Human	perox	632	Α					
IDVWLGGLALNFLPY	1468	13	ruman	thyroid perox	032	А					
IDVWLGGLASNFLPY	1400	15	Human	thyroid	632	Α					
ID THE COLLEGE AND A	1469			perox							
IDVWLGGLAKNFLPY		15	Human	thyroid	632	Α					
	<u>1470</u>			perox							
IDVWLGGLADNFLPY	1471	15	Human	thyroid	632	Α					
IDVYLGGLAENFLPY	<u>1471</u>	15	Human	perox thyroid	632	Α					
IDVI LOGEALINI EL I	1472	13	ruman	perox	032	11					
IDVLLGGLAENFLPY		15	Human	thyroid	632	Α					
	1473			perox							
IDVSLGGLAENFLPY		15	Human	thyroid	632	Α					
IDVKLGGLAENFLPY	<u>1474</u>	15	Human	perox thyroid	632	Α					
IDVKLUGLAENFLFI	1475	13	riuman	perox	032	Α.					
IDVDLGGLAENFLPY	1175	15	Human	thyroid	632	Α					
	1476			perox							
IDVWLGGLAENYLPY		15	Human	thyroid	632	Α					
IDIANI GGI ADAMAN	<u>1477</u>	1.5	Human	perox	632						
IDVWLGGLAENVLPY	1478	15	Human	thyroid perox	032	Α					
IDVWLGGLAENSLPY	1470	15	Human	thyroid	632	Α					
	1479			perox							
IDVWLGGLAENKLPY		15	Human	thyroid	632	Α					
	<u>1480</u>		**	perox	620						
IDVWLGGLAENDLPY	1481	15	Human	thyroid perox	632	Α					
IYVWLGGLAENFLPY	1401	15	Human	thyroid	632	Α					
TT TO SET LET T	1482		***********	perox							
ILVWLGGLAENFLPY		15	Human	thyroid	632	Α					
	<u>1483</u>			perox							
ISVWLGGLAENFLPY	1484	15	Human	thyroid perox	632	Α					
IKVWLGGLAENFLPY	1464	15	Human	thyroid	632	Α					
IK V W EGGENERATES T	1485			perox	052	••					
IEVWLGGLAENFLPY		15	Human	thyroid	632	Α					
	1486			perox							
IDVWLGGLAENFLPF	1.407	15	Human	thyroid perox	632	Α					
IDVWLGGLAENFLPL	<u>1487</u>	15	Human	thyroid	632	Α					
ID LOGENERI EFE	1488			perox	002	••					
IDVWLGGLAENFLPS		15	Human	thyroid	632	Α					
	1489			perox							
IDVWLGGLAENFLPK	1490	15	Human	thyroid perox	632	Α					
IDVWLGGLAENFLPD	1490 1491	15	Human	thyroid	632	Α					
	1471			,							

HLA-DQ SUPERTYPES									
	SEQ ID								
Sequence	NO.	AA	Organism	Protein	Position	Analog			
				perox					
IDVWLGGLAENFYPY		15	Human	thyroid	632	Α			
IDVWLGGLAENFVPY	<u>1492</u>	15	Human	perox thyroid	632	Α			
IDV WLGGLAENFVF I	1493	13	riuman	perox	032	Λ.			
IDVWLGGLAENFSPY		15	Human	thyroid	632	Α			
	<u>1494</u>		**	perox	632				
IDVWLGGLAENFKPY	1495	15	Human	thyroid perox	032	Α			
IDVWLGGLAENFDPY	1422	15	Human	thyroid	632	Α			
	1496			perox					
IDVWLGGLAEYFLPY	1497	15	Human	thyroid perox	632	Α			
IDVWLGGLAELFLPY	1497	15	Human	thyroid	632	Α			
	1498			perox					
IDVWLGGLAESFLPY		15	Human	thyroid	632	Α			
IDVWLGGLAEKFLPY	<u>1499</u>	15	Human	perox thyroid	632	Α			
IDV WEGGEAERI'EI I	1500	13	Haman	perox	032	**			
IDVWLGGLAEDFLPY		15	Human	thyroid	632	Α			
IDAMA CCI A POPI DV	1501	15	Human	perox	632	Α			
IDVWLGGLAEQFLPY	1502	13	ruman	thyroid perox	032	A			
IDVWLGGLYENFLPY		15	Human	thyroid	632	Α			
	1503			perox					
IDVWLGGLLENFLPY	1504	15	Human	thyroid perox	632	Α			
IDVWLGGLSENFLPY	1304	15	Human	thyroid	632	Α			
	1505			perox					
IDVWLGGLKENFLPY	1506	15	Human	thyroid perox	632	Α			
IDVWLGGLDENFLPY	1300	15	Human	thyroid	632	Α			
	1507			perox					
IDVWLGGYAENFLPY		15	Human	thyroid	632	Α			
IDVWLGGVAENFLPY	<u>1508</u>	15	Human	perox thyroid	632	Α			
ID V W LOG V ALIVI LI I	1509	13	Trumum	perox	032	**			
IDVWLGGSAENFLPY		15	Human	thyroid	632	Α			
IDVWLGGKAENFLPY	<u>1510</u>	15	Human	perox thyroid	632	Α			
ID V W EGGKAENTEI T	1511	13	man	perox	032	**			
IDVWLGGDAENFLPY		15	Human	thyroid	632	Α			
IDVWLGYLAENFLPY	1512	15	Human	perox thyroid	632	Α			
IDVWLGILAENFLFI	1513	13	ruman	perox	032	Α			
IDVWLGLLAENFLPY		15	Human	thyroid	632	Α			
IDIUM COL LEVEL DI	<u>1514</u>	1.5	TT	perox	632	Α			
IDVWLGSLAENFLPY	1515	15	Human	thyroid perox	032	А			
IDVWLGKLAENFLPY	1510	15	Human	thyroid	632	Α			
	<u>1516</u>			perox	600				
IDVWLGDLAENFLPY	1517	15	Human	thyroid perox	632	Α			
IDVWLYGLAENFLPY	1311	15	Human	thyroid	632	Α			
	<u>1518</u>			perox					

		ILA-DQ	SUPERTYPES			
	SEQ ID					
Sequence	NO.	AA	Organism	Protein	Position	Analog
IDVWLLGLAENFLPY		15	Human	thyroid	632	Α
IDVWLSGLAENFLPY	<u>1519</u>	15	Human	perox thyroid	632	Α
ID V W L3GLAENFLF I	1520	13	Human	perox	032	Α
IDVWLKGLAENFLPY		15	Human	thyroid	632	Α
IDVWLDGLAENFLPY	<u>1521</u>	15	Human	perox thyroid	632	Α
IDV W LDGLAENFLF I	1522	13	numan	perox	032	Λ
IDVWYGGLAENFLPY		15	Human	thyroid	632	Α
TOURING OF A EXIET DV	<u>1523</u>	1.5	********	perox	632	
IDVWVGGLAENFLPY	1524	15	Human	thyroid perox	632	Α
IDVWSGGLAENFLPY	1021	15	Human	thyroid	632	Α
	1525			perox		
IDVWKGGLAENFLPY	1526	15	Human	thyroid perox	632	Α
IDVWDGGLAENFLPY	1320	15	Human	thyroid	632	Α
	<u>1527</u>			perox		
IDYWLGGLAENFLPY	1500	15	Human	thyroid	632	Α
IDLWLGGLAENFLPY	<u>1528</u>	15	Human	perox thyroid	632	Α
	1529			perox		
IDSWLGGLAENFLPY	4500	15	Human	thyroid	632	Α
IDKWLGGLAENFLPY	<u>1530</u>	15	Human	perox thyroid	632	Α
IDK WEGGENEIGI EN	1531		114111411	perox	052	••
IDDWLGGLAENFLPY		15	Human	thyroid	632	Α
IDVWLGGLAENFLYY	<u>1532</u>	15	Human	perox thyroid	632	Α
IDV WEGGLAENFE I	1533	13	Human	perox	032	Λ
IDVWLGGLAENFLLY		15	Human	thyroid	632	Α
IDVWLGGLAENFLSY	<u>1534</u>	15	Human	perox thyroid	632	Α
IDV WLGGLAENFLS I	1535	13	пишап	perox	032	A
IDVWLGGLAENFLKY		15	Human	thyroid	632	Α
TOTAL COLUMNIA	<u>1536</u>		**	perox	620	
IDVWLGGLAENFLDY	1537	15	Human	thyroid perox	632	Α
YDVWLGGLAENFLPY	1337	15	Human	thyroid	632	Α
	<u>1538</u>			perox		
LDVWLGGLAENFLPY	1539	15	Human	thyroid perox	632	Α
SDVWLGGLAENFLPY	1335	15	Human	thyroid	632	Α
	1540			perox		
KDVWLGGLAENFLPY	1541	15	Human	thyroid	632	Α
DDVWLGGLAENFLPY	<u>1541</u>	15	Human	perox thyroid	632	Α
	1542			perox		

TABLE 24

Sequence NO. 1 2 1 AAAKAAAAAAYAA 1429 424 (44)YAAAAAKAAA 1430 26 AAFAAAKTAAAFA 1431 49 YAAFAAAKTAAAFA 1432 36 YAAFAAAKTAAAFA 1433 39 AHAAHAAHAAHAAHAA 1435 10932 309 5389 VPDTKVNFYAWKRME 1436 730 >46666.67 >147058.82 WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 144	H	LA-DQ SU	PERTYPES		
AAAKAAAAAYAA 1429 424 (44)YAAAAAAKAAA 1430 26 AAFAAAKTAAAFA 1431 49 YAAFAAAKTAAAFA 1432 36 YAAFAAAKTAAAFA 1433 39 AHAAHAAHAAHAAHAA 1434 58 VLERYLLEAKEAENI 1435 10932 309 5389 VPDTKVNFYAWKRME 1436 730 >46666.67 >147058.82 WKRMEVGQQAVEWWQ 1437 136666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEWWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 EAENITTGTAEHTSL 1447 316 8300 EAENITTGTAEHTSL 1449 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLLKDLEEG 1452 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKDLEEG 1452 89285.71 502 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SCHORNFDIPEEIK 1454 19365 208 774 QLQPQKEDAAVTIY 1455 26205 579 2145 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVEU 1469 1553 8413 359 GSHLVEALYLVCGER 1463 >89285.71 2491 677 GSBUVEALYLVCGER 1463 89285.71 2491 677 GSBUVEALYLVCGER 1463 89285.71 2491 677 GSBUVEALYLVCGER 1463 89285.71 2491 677 GSBUVEGLANFLPY 1466 204 138 13 IDVWLGGLANFLPY 1468 49 457 52 IDVWLGGLANFLPY 1468 175 1251 40 IDVWLGGLANFLPY 1466 175 1251 40			DQB1*030	DQB1*030	DQB1*020
(44)YAAAAAAKAAA 1430 26 AAFAAAKTAAAFA 1431 49 YAAFAAAKTAAAFA 1432 36 YAAFAAAKTAAAFA 1433 39 AHAAHAAHAAHAA 1434 58 VLERYLLEAKEAENI 1435 10932 309 5389 VPDTKVNFYAWKRME 1436 730 >46666.67 >147058.82 WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGIRSL 1443 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 11766 RLFDNASLRAHRLHQ </th <th>Sequence</th> <th>NO.</th> <th>11</th> <th>2</th> <th>1</th>	Sequence	NO.	11	2	1
AÁFAAAKTAAAFA YAAFAAAKTAAAFA YAAFAAAKTAAAFA YAAFAAAKTAAAFA 1432 36 YAAFAAAKTAAAFA 1433 39 AHAAHAAHAAHAAHAA 1434 58 VLERYLLEAKEAENI 1435 10932 309 5389 VPDTKVNFYAWKRME 1436 730 >46666.67 147058.82 WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEWWQGLALSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 A6666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 36206.9 11766 QLAFDTYQEFEEAYI 1449 898285.71 673 35 ISLLLIQSWLEPVQF 1450 1852 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLLKDLEEG 1452 889285.71 673 35 ISLLLIQSWLEPVQF 1450 1451 14164 8337 731 SDSNYDLLKDLEEG 1452 889285.71 673 35 ISLLLIQSWLEPVQF 1450 1451 14164 8337 731 SDSNYDLLKDLEEG 1452 1454 19365 208 774 QLQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 49285.71 806 VEALYLVCGER 1463 889285.71 806 VEALYLVCGER 1463 889285.71 806 VEALYLVCGER 1463 89285.71 806 VEALYLVCGER 1464 27334 514 VEALYLVTGERGFFY 1465 10VWLGGLASNFLPY 1468 10VWLGGLASNFLPY 1469 175 1251 10VWLGGLASNFLPY 1469 175 1100 10247 84166.67 1100 10247 84166.67	AAAKAAAAAYAA	1429	424		
YAAFAAAKTAAAFA 1432 36 YAAFAAAKTAAAFA 1433 39 AHAAHAAHAAHAA 1434 58 VPDTKVNFYAWKRME 1436 730 >46666.67 >147058.82 WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RAFDARSHAHRLHQ 1448 996 7287 PDASALRAHRLHQ 1448 996 733 35 </td <td>(44)YAAAAAKAAA</td> <td>1430</td> <td>26</td> <td></td> <td></td>	(44)YAAAAAKAAA	1430	26		
YAAFAAAKTAAAFA 1433 39 AHAAHAAHAAHAAHAA 1434 58 VLERYLLEAKEAENI 1435 10932 309 5389 VPDTKVNFYAWKRME 1436 730 >46666.67 >147058.82 WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHYDKAVSGIRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITATOFFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 11766 OLAFDTYGEFEA	AAFAAAKTAAAFA	1431	49		
AHAAHAAHAAHAA AHAAHAAHAAHAA VLERYLLEAKEAENI VLERYLLEAKEAENI VPDTKVNFYAWKRME 435 WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 A6666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ QLAFDTYQEFEEAYI 1449 8996 36206.9 11766 QLAFDTYQEFEEAYI 1449 8996 36206.9 11766 YSSUNYDLLKDLEEG 1451 SDSNVYDLLKDLEEG 1452 89285.71 4136 503 KIFGSLAFLPESFDGDPA CLKDRRNFDIPEEIK QLQFQKEDAAVTIY QKEDAAVTIYEMLQN 1455 26205 579 2145 CKEDSHCAWTIVEVEL 1458 708 708 708 709 709 709 709 70	YAAFAAAKTAAAFA	1432	36		
VLERYLLEAKEAENI 1435 10932 309 5389 VPDTKVNFYAWKRME 1436 730 >466666.67 >147058.82 WKRMEVGQQAVEVWQ 1437 13666 12146 139 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 <	YAAFAAAKTAAAFA	1433	39		
VPDTKVNFYAWKRME 1436 730 >46666.67 >147058.82 WKRMEVGQQAVEWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3699 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 7287 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 98285.71 673 35 ISLLIQSWLEPVQF 1450 >892	АНААНААНААНАА	1434	58		
WKRMEVGQQAVEVWQ 1437 13666 12146 159 VGQQAVEVWQGLALL 1438 1807 4407 838 VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHYDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGAEHTSL 1447 316 8300 11766 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 ×89285.71 4136 503 KIFGSLAFLPESEDGDPA 1453 320<	VLERYLLEAKEAENI	1435	10932	309	5389
VGQQAVEVWQGLALL VGQQAVEVWQGLALL VGQQAVEVWQGLALL VGQQAVEVWQGLALL VGQQAVEVWQGLALLSEAVL VEVWQGLALLSEAVL VGQAVEVWQGLALLSEAVL VGQAVEVWQGLALLSEAVL VGGAVEVWQGLALLSEAVL VGAVSGLRSL VGAVEVWGGALLVNSSQPWEP VGAVAVAVA VGAVAVAVA VGAVAVAVAVA VGAVAVAVAV	VPDTKVNFYAWKRME	1436	730	>46666.67	>147058.82
VEVWQGLALLSEAVL 1439 19 14 98 GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 74<	WKRMEVGQQAVEVWQ	1437	13666	12146	159
GLALLSEAVLRGQAL 1440 107 16963 6742 SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13365 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLAFTTY 1447 316 8300 RLAFTY 1449 SERVEST 1673 35 155LLLQSWLEPVQF 1450 >89285.71 673 35 15LLLQSWLEPVQF 1450 >89285.71 673 35 15LLLQSWLEPVQF 1450 >89285.71 673 35 15LLLQSWLEPVQF 1450 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQPGKEDAAVTIY 1455 26205 579 2145 QLQPGKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVEVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 QHLCGSHLVEALYLVCGER 1463 89285.71 2491 677 GSDLVEALYLVCGER 1463 89285.71 2491 677 GSDLVEALYLVCGER 1463 89285.71 2491 677 GSDLVEALYLVCGER 1463 89285.71 806 VEALYLVCGERGFLY 1466 204 138 13 IDVWLGGLASNFLPY 1467 85 358 63 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLASNFLPY 1461 170 10247 >4166.67 IDVWLGGLASNFLPY 1471 296 1762 12	VGQQAVEVWQGLALL	1438	1807	4407	838
SEAVLRGQALLVNSS 1441 55 36395 9755 RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 11766 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNYDLLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 <	VEVWQGLALLSEAVL	1439	19	14	98
RGQALLVNSSQPWEP 1442 302 14393 13362 LQLHVDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVNVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 QHLCGSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 89285.71 2491 677 GSDLVEALYLVCGER 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLANFLPY 1468 49 457 52 IDVWLGGLANFLPY 1468 49 457 52 IDVWLGGLANFLPY 1469 175 1251 40 IDVWLGGLANFLPY 1469 175 1251 40 IDVWLGGLANFLPY 1461 170 10247 >4166.67 IDVWLGGLANFLPY 1461 170 10247 >4166.67	GLALLSEAVLRGQAL	1440	107	16963	6742
LQLHYDKAVSGLRSL 1443 88 7842 7590 KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 11766 QLAFDTYQEFEEAYI 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 662 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 14164 8337 731 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQPGKEDAAVTIY 1455 26205 579 2145 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVEVEI	SEAVLRGQALLVNSS	1441	55	36395	9755
KEAISPPDAASAAPL 1444 458 960 7287 PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 11766 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNYDLLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 774 14136 503 CLKDRRNFDIPEEIK 1454 19365 208 774 1456 515 153 1685 STGWNETIVENLLAN 1456 515 153 1685 1685 579 2145 2123 465 465 465 465 465 465 465 465	RGOALLVNSSOPWEP	1442	302	14393	13362
PPDAASAAPLRTITA 1445 20 3869 3631 SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLLQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 ×92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 889285.71 2491 677 GSDLVEALYLVCGER 1462 S89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1468 49 457 52 IDVWLGGLAENFLPY 1468 49 457 52 IDVWLGGLAENFLPY 1468 49 175 1251 40 IDVWLGGLAENFLPY 1468 49 175 1251 40 IDVWLGGLAENFLPY 1469 175 1251 40 IDVWLGGLAENFLPY 1469 175 1251 40 IDVWLGGLAENFLPY 1469 177 110247 >4166.67	LOLHVDKAVSGLRSL	1443	88	7842	7590
SAAPLRTITADTFRK 1446 301 >46666.67 1100 EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 774<	KEAISPPDAASAAPL	1444	458	960	7287
EAENITTGTAEHTSL 1447 316 8300 RLFDNASLRAHRLHQ 1448 996 >36206.9 11766 QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPVQF 1450 >89285.71 562 5234 NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 20 20 774 QKEDAAVTIYEMLQN 1456 515 153 1685 57GWNETIVENLLAN 1457 47081 5041 322 22 2145 20 205 579 2145 2145 20 205 579 2145 2145 2145 2145 22 22 22 22 22 22 22 22 22 22 22 22	PPDAASAAPLRTITA	1445	20	3869	3631
RLFDNASLRAHRLHQ RLAFDTYQEFEEAYI RLEDNASLRAHRLHQ RLAFDTYQEFEEAYI RSULYGASDSNVYDL RSULYGASDSNVYLG RSULYGASDSNV RSULYGASDSNVYDL R	SAAPLRTITADTFRK	1446	301	>46666.67	1100
QLAFDTYQEFEEAYI 1449 >89285.71 673 35 ISLLLIQSWLEPYQF 1450 >89285.71 562 5234 NSLVYGASDSNYYDL 1451 14164 8337 731 SDSNYYDLLKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY	EAENITTGTAEHTSL	1447	316	8300	
ISLLLIQSWLEPVQF	RLFDNASLRAHRLHQ	1448	996	>36206.9	11766
NSLVYGASDSNVYDL 1451 14164 8337 731 SDSNVYDLKKDLEEG 1452 >89285.71 4136 503 KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETTVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLASNFLPY 1466 204 138 13 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1469 175 1251 40 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1469 170 10247 >4166.67	QLAFDTYQEFEEAYI	1449	>89285.71	673	35
SDSNVYDLLKDLEEG	ISLLLIOSWLEPVOF	1450	>89285.71	562	5234
KIFGSLAFLPESFDGDPA 1453 320 CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1459 47081 5041 322 ETIVENLLANVYHQR 1459 4702 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 889285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLALNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLALNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1461 170 10247 >4166.67	NSLVYGASDSNVYDL	1451	14164	8337	731
CLKDRRNFDIPEEIK 1454 19365 208 774 QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 465 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSBLVEALYLVCGER 1463 >89285.71 2491 677 GSDLVEALYLVCGERGFFY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLASPILPY 1466 204 138 13 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40	SDSNVYDLLKDLEEG	1452	>89285.71	4136	503
QLQQFQKEDAAVTIY 1455 26205 579 2145 QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 2401 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 806 77 GSDLVEALYLVCGER 1463 >89285.71 806 77 VEALYLVTGERGFFY 1464 27334 514 514 VEALYLVTGERGFFY 1465 20021 564 110 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLALNFLPY 1468 49 457 52 IDVWLGGLALNFLPY 1468 49 457 52 IDVWLGGLAKNFLPY 1469 175	KIFGSLAFLPESFDGDPA	1453	320		
QKEDAAVTIYEMLQN 1456 515 153 1685 STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 724 904	CLKDRRNFDIPEEIK	1454	19365	208	774
STGWNETIVENLLAN 1457 47081 5041 322 ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 OHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLANFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1470 170 10247 >4166.67 IDVWLG	QLQQFQKEDAAVTIY	1455	26205	579	2145
ETIVENLLANVYHQR 1458 >92592.59 >75000 344 KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 9 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGERGFLY 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1471 296 1762 12 <td>QKEDAAVTIYEMLQN</td> <td>1456</td> <td>515</td> <td>153</td> <td>1685</td>	QKEDAAVTIYEMLQN	1456	515	153	1685
KEDSHCAWTIVRVEI 1459 4102 2123 465 MSYNLLGFLQRSSNT 1460 724 >51219.51 359 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVTGERGFFY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 DVULGGLAENFLPY 1466 204 138 13 13 IDVWLGGLAYNFLPY 1467 85 358 63 63 IDVWLGGLALNFLPY 1468 49 457 52 10VWLGGLASNFLPY 1469 175 1251 40 10VWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLADNFLPY 1471 296 1762 12 12	STGWNETIVENLLAN	1457	47081	5041	322
MSYNLLGFLQRSSNT 1460 724 >51219.51 QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLALNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1470 170 10247 >4166.67	ETIVENLLANVYHQR	1458	>92592.59	>75000	344
QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLASNFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1471 296 1762 12	KEDSHCAWTIVRVEI	1459	4102	2123	465
QHLCGSHLVEALYLV 1461 2553 8413 359 GSHLVEALYLVCGER 1462 >89285.71 2491 677 GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLASNFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1471 296 1762 12	MSYNLLGFLQRSSNT	1460	724	>51219.51	
GSDLVEALYLVCGER 1463 >89285.71 806 VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLASNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLANFLPY 1471 296 1762 12	OHLCGSHLVEALYLV	1461	2553	8413	359
VEALYLVCGERGFLY 1464 27334 514 VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLALNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLADNFLPY 1471 296 1762 12	GSHLVEALYLVCGER	1462	>89285.71	2491	677
VEALYLVTGERGFFY 1465 20021 564 IDVWLGGLAENFLPY 1466 204 138 13 IDVWLGGLAYNFLPY 1467 85 358 63 IDVWLGGLAYNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLADNFLPY 1471 296 1762 12	GSDLVEALYLVCGER	1463	>89285.71	806	
IDVWLGGLAENFLPY	VEALYLVCGERGFLY	1464	27334	514	
IDVWLGGLAYNFLPY	VEALYLVTGERGFFY		20021	564	
IDVWLGGLALNFLPY	IDVWLGGLAENFLPY	1466	204	138	13
IDVWLGGLALNFLPY 1468 49 457 52 IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLADNFLPY 1471 296 1762 12	IDVWLGGLAYNFLPY		85	358	63
IDVWLGGLASNFLPY 1469 175 1251 40 IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLADNFLPY 1471 296 1762 12	IDVWLGGLALNFLPY	_	49	457	52
IDVWLGGLAKNFLPY 1470 170 10247 >4166.67 IDVWLGGLADNFLPY 1471 296 1762 12	IDVWLGGLASNFLPY		175	1251	40
IDVWLGGLADNFLPY <u>1471</u> 296 1762 12	IDVWLGGLAKNFLPY	_	170	10247	>4166.67
	IDVWLGGLADNFLPY		296	1762	
10 14/2 101 100 30	IDVYLGGLAENFLPY	1472	161	186	30

HLA-DQ SUPERTYPES									
	SEQ ID	DQB1*030	DQB1*030	DQB1*020					
Sequence	NO.	1	2	1					
IDVLLGGLAENFLPY	1473	166	437	27					
IDVSLGGLAENFLPY	1474	188	277	48					
IDVKLGGLAENFLPY	1475	724	5511	41					
IDVDLGGLAENFLPY	1476	218	73	17					
IDVWLGGLAENYLPY	1477	223	110	19					
IDVWLGGLAENVLPY	1478	84	82	15					
IDVWLGGLAENSLPY	1479	116	125	25					
IDVWLGGLAENKLPY	1480	353	5189	51					
IDVWLGGLAENDLPY	1481	240	60	22					
IYVWLGGLAENFLPY	1482	170	237	13					
ILVWLGGLAENFLPY	1483	216	147	10.0					
ISVWLGGLAENFLPY	1484	132	286	18					
IKVWLGGLAENFLPY	1485	180	220	37					
IEVWLGGLAENFLPY	1486	158	145	23					
IDVWLGGLAENFLPF	1487	111	177	3.6					
IDVWLGGLAENFLPL	1488	182	114	17					
IDVWLGGLAENFLPS	1489	134	249	27					
IDVWLGGLAENFLPK	1490	261	231	23					
IDVWLGGLAENFLPD	1490	115	91	20					
IDVWLGGLAENFYPY	1491	324	203	37					
IDVWLGGLAENFVPY	1492	346	272	12					
IDVWLGGLAENFSPY	1493 1494	131	193	47					
IDVWLGGLAENFKPY	1494 1495	195	262	310					
IDVWLGGLAENFDPY	1495	364	90	32					
IDVWLGGLAEYFLPY	1496 1497	151	88	14					
IDVWLGGLAEIFLFI	1497	107	81	22					
IDVWLGGLAESFLPY		60	64	49					
IDVWLGGLAESFLFT	<u>1499</u>	68	112	66					
IDVWLGGLAEDFLPY	<u>1500</u>	357	120	23					
	<u>1501</u>	357 167	120	2.5 9.7					
IDVWLGGLAEQFLPY	<u>1502</u>	912	697	6.4					
IDVWLGGLYENFLPY	<u>1503</u>		1734	58					
IDVWLGGLLENFLPY	<u>1504</u>	810 242	1734	38 37					
IDVWLGGLSENFLPY	1505								
IDVWLGGLKENFLPY	<u>1506</u>	15907	>2800	25					
IDVWLGGLDENFLPY	<u>1507</u>	>19230.77	637	18					
IDVWLGGYAENFLPY	<u>1508</u>	900	492	39					
IDVWLGGVAENFLPY	<u>1509</u>	982	327	75					
IDVWLGGSAENFLPY	<u>1510</u>	427	755	166					
IDVWLGGKAENFLPY	<u>1511</u>	517	633	398					
IDVWLGGDAENFLPY	<u>1512</u>	11114	2074	11					
IDVWLGYLAENFLPY	<u>1513</u>	15215	1121	31					
IDVWLGLLAENFLPY	<u>1514</u>	2986	180	39					
IDVWLGSLAENFLPY	<u>1515</u>	654	278	72					
IDVWLGKLAENFLPY	<u>1516</u>	2333	20023	81					
IDVWLGDLAENFLPY	<u>1517</u>	>44642.86	370	18					
IDVWLYGLAENFLPY	<u>1518</u>	2171	442	18					

	HLA-DQ SU	PERTYPES		
-	SEQ ID	DQB1*030	DQB1*030	DQB1*020
Sequence	NO.	1	2	1
IDVWLLGLAENFLPY	1519	4903	455	47
IDVWLSGLAENFLPY	1520	3043	373	98
IDVWLKGLAENFLPY	1521	41667	1115	55
IDVWLDGLAENFLPY	1522	13325	357	43
IDVWYGGLAENFLPY	1523	375	224	43
IDVWVGGLAENFLPY	1524	128	158	14
IDVWSGGLAENFLPY	1525	451	128	15
IDVWKGGLAENFLPY	1526	256	346	41
IDVWDGGLAENFLPY	1527	2086	299	112
IDYWLGGLAENFLPY	1528	503	342	49
IDLWLGGLAENFLPY	1529	1292	661	25
IDSWLGGLAENFLPY	<u>1530</u>	508	276	35
IDKWLGGLAENFLPY	<u>1531</u>	579	534	62
IDDWLGGLAENFLPY	1532	219	101	85
IDVWLGGLAENFLYY	1533	341	387	154
IDVWLGGLAENFLLY	1534	649	491	52
IDVWLGGLAENFLSY	1535	425	676	54
IDVWLGGLAENFLKY	1536	2266	995	111
IDVWLGGLAENFLDY	1537	371	149	49
YDVWLGGLAENFLPY	1538	482	214	59
LDVWLGGLAENFLPY	1539	180	216	29
SDVWLGGLAENFLPY	1540	154	232	19
KDVWLGGLAENFLPY	1541	348	254	54
DDVWLGGLAENFLPY	1542	241	158	48

TABLE 25

		HLA-	DR SUPERTYPE			
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
AC-		18	A2	MHC derived	Unknown	Anaiog
NPTKHKWEAAHVAE						
QLAA	1543					
DDYVKQYTKQYTKQ		19	Artificial			
NTLKK	<u>1544</u>		sequence			
AAAKAAAAAAYAA		13	Artificial			Α
	<u>1545</u>		sequence			Α
AC-	1546	13	Artificial			А
AAAKAAAAAYAA (20)AYA(20)A(20)A(20)	1340	13	sequence Artificial			Α
K(20)A(20)	1547	13	sequence			- ~
AC-	1011	13	Artificial			Α
AAAKATAAAAYAA	1548		sequence			
AC-		13	Artificial			Α
AAAKAAAAAAFAA	<u>1549</u>		sequence			
AC-		13	Artificial			Α
AAAKATAAAA(10)AA	<u>1550</u>		sequence			
AC-		13	Artificial			Α
AAAKATAAAA(23)AA	<u>1551</u>		sequence			
AAKAAAAAAA(10)AA		13	Artificial			Α
	1552		sequence			
AAYAAAATAKAAA		13	Artificial			Α
	<u>1553</u>	12	sequence			Α
AALAAAAAKAAA	1554	13	Artificial			А
AAEAAAATAKAAA	1334	13	sequence Artificial			Α
AABAAAATAKAAA	1555	13	sequence			- 1
AAYJJAAAAKAAA	1000	13	Artificial			Α
AATSAAAAAA	1556		sequence			•••
AAYAAAAJJKAAA		13	Artificial			Α
	1557		sequence			
AFLRAAAAAAFAA		13	Artificial			Α
	<u>1558</u>		sequence			
AFLRQAAAAAFAAY		14	Artificial			Α
	<u>1559</u>		sequence			
AAFAAAKTAAAFA	1560	13	Artificial			Α
W E W.T E.	<u>1560</u>		sequence			Α
YAAFAAAKTAAAFA	1561	14	Artificial			А
AALKATAAAAAA	<u>1561</u>	13	sequence Artificial			Α
AALKATAAAAAA	1562	13	sequence			^
YAR(15)ASQTTLKAKT	1502	14	Artificial			
(15)110@112111111	1563	• •	sequence			
YARF(33)QTTLKAKT		14	Artificial			
	1564		sequence			
PKYFKQRILKFAT		13	Artificial			Α
	1565		sequence			
PKYFKQGFLKGAT		13	Artificial			Α
	<u>1566</u>		sequence			
PKYGKQIDLKGAT	1000	13	Artificial			Α
4 4 PPPPPCCCCC 4	<u>1567</u>	12	sequence			
AAFFFFFGGGGGA	1568	13	Artificial			
AADFFFFFFFDA	1508	13	sequence Artificial			
AADETTTTTDA	1569	13	sequence			
AAKGIKIGFGIFA	1000	13	Artificial			
	1570		sequence			
AAFIFIGGGKIKA		13	Artificial			
	1571		sequence			
AAKIFIGFFIDGA		13	Artificial			
	1572		sequence			

		HLA-	DR SUPERTY	PE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
AAFIGFGKIKFIA		13	Artificial			
	<u>1573</u>		sequence			
AAKIGFGIKIGFA		13	Artificial			
+ + PKIOKECIEE+	<u>1574</u>	12	sequence			
AAFKIGKFGIFFA	1575	13	Artificial sequence			
AADDDDDDDDDDA	15/5	13	Artificial			
	<u>1576</u>		sequence			
(43)AAIGFFFFKKGIA		14	Artificial			
(43) A A EPOLEKICKEA	<u>1577</u>	14	sequence			
(43)AAFFGIFKIGKFA	1578	14	Artificial sequence			
(43)AADFGIFIDFIIA	10.70	14	Artificial			
,	<u>1579</u>		sequence			
(43)AAIGGIFIFKKDA		14	Artificial			
(43) A A PICECKIKEI A	<u>1580</u>	13	sequence			
(43)AAFIGFGKIKFIA	1581	13	Artificial sequence			
(43)AAKIGFGIKIGFA	1501	13	Artificial			
	<u>1582</u>		sequence			
(43)AAFKIGKFGIFFA		13	Artificial			
	<u>1583</u>		sequence			
AAAKAAAAAAAAF	1584	13	Artificial sequence			
AAAKAAAAAAAFA	1304	13	Artificial			
	<u>1585</u>		sequence			
AAAKAAAAAAFAA		13	Artificial			
	<u>1586</u>	13	sequence			
AAAKAAAAFAAAA	1587	13	Artificial sequence			
FAAAAAAAAAAA	1507	13	Artificial			
	<u>1588</u>		sequence			
AAAAAAAAAAAN		13	Artificial			
AAAAAAAAAAANA	<u>1589</u>	13	sequence Artificial			
AAAAAAAAAAA	1590	13	sequence			
AAANAAAAAAAA	1370	13	Artificial			
	<u>1591</u>		sequence			
AAAAAAAAAAA	1500	13	Artificial			
AAAAAAAAAAA	<u>1592</u>	13	sequence			
AAAAASAAAAAA	1593	13	Artificial sequence			
ASAAAAAAAAAA		13	Artificial			
	<u>1594</u>		sequence			
AFAAAKTAA	1505	9	Artificial			
YARFLALTTLRARA	<u>1595</u>	14	sequence Artificial			Α
TAKI LALI ILKAKA	1596	1-4	sequence			^
YAR(15A)SQTTLKAKT		14	Artificial			Α
	<u>1597</u>		sequence			
YAR(15A)RQTTLKAA	1.500	14	Artificial			Α
A (15A)RQTTLKAAA	<u>1598</u>	11	sequence Artificial			Α
(154)KGI IPKAKA	1599		sequence			Α.
(16A)RQTTLKAAA		11	Artificial			Α
	1600		sequence			
(46)AAKTAAAFA	1601	10	Artificial			
(39)AAAATKAAA	1601	10	sequence Artificial			
(22)MANATIKANA	1602		sequence			
(52)AAAATKAAAA		11	Artificial			
	<u>1603</u>		sequence			

		HLA-	DR SUPERTYPE	3		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
(55)AAAATKAAAA		11	Artificial		- "	
	1604		sequence			
A(14)AAAKTAAA	1605	10	Artificial			
AA(14)A(35)ATKAAA	1003	12	sequence Artificial			
A	1606		sequence			
AA(14)AA(36)TKAAA		12	Artificial			
A	<u>1607</u>	10	sequence Artificial			
AFAAAKTAA(72)	1608	10	sequence			
(49)AAAKT(64)AAA		10	Artificial			
	1609		sequence			
(49)AAAKTA(64)AA	1610	10	Artificial sequence			
HOAISPRTLNGPGPGS	1010	20	Artificial			
PAIF	1611		sequence			
YAAFAAAKTAAAFA		14	Artificial			
TEGRCLHYTVDKSKP	<u>1612</u>	16	sequence Bee Venom		103	
K	1613	10	Bee venom		103	
AWVAWRNRCK	1614	0	Chicken	HEL	107	
IVSDGNGMNAWVAW		18	Chicken	HEL	98	
RNRC	<u>1615</u>					
PHHTALRQAILSWGE	1616	20	DPw4 binder			
LMTLA WMYYHGQRHSDEHH	<u>1616</u>	15	EBV	LMP	183	
Н	1617		LD.			
YIVMSDWTGGA	1618	15	EBV	LMP	41	
AHAAHAAHAAHA		16	HA			Α
AA	<u> 1619</u>	2.5	нву	core	1	
MDIDPYKEFGATVEL LSFLPSDFFP	1620	25	поч	core	1	
GMLPVCPLIPGSSTTS	1311	19	HBV	env	102	
TGP	1621					
LGFFPDHQLDPAFRA NT	1622	17	HBV	env	11	
GYKVLVLNPSV	1623	11	HCV	NS3	1248	
LMAFTAAVTS		10	HCV	NS4	1790	
TFALWRVSAEEY	1624	12	HCV	NS5	2079	
ALWRVSAEEY	1625	10	HCV	NS5	2081	
	1626	13	HCV	NS5	2088	
EEYVEIRQVGDFH	<u>1627</u>			NSS	2088	
VGGVYLLPRRGPRLG V	1628	16	HCV			
VGGAYLLPRRGPRLG	1020	16	HCV			Α
V	1629					
VGGVALLPRRGPRLG	1630	16	HCV			Α
V VGGVYALPRRGPRLG	1030	16	HCV			Α
V	1631					
VGGVYLAPRRGPRLG	1622	16	HCV			Α
V VGGVYLLARRGPRLG	<u>1632</u>	16	HCV			Α
V	1633	10	iic v			^
VGGVYLLPARGPRLG		16	HCV			Α
V	<u>1634</u>	16	UCV			
VGGVYLLRRAGPRLG V	1635	16	HCV			Α
GAPLGGAARALAHGV		15	HCV			
GAALGGAARALAHG		15	HCV			Α
V	1637					

		HLA-	DR SUPERTY	PE		
	SEQ ID NO.					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
GAPLAGAARALAHGV	<u>1638</u>	15	HCV			Α
GAPLGAAARALAHGV	1639	15	HCV			Α
GAPLGGLARALAHGV	1640	15	HCV			Α
GAPLGGALRALAHGV	1641	15	HCV			Α
GAPLGGAAAALAHG V	1642	15	HCV			Α
GAPLGGAARLLAHGV	1643	15	HCV			Α
GAPLGGAARAAAHG		15	HCV			Α
V	<u>1644</u>					
GAPLGGAARALAAGV	1645	15	HCV			Α
FPDWQNYTPGPGTRF	1646	15	HIV	NEF	200	
RFPLTFGWCFKLVPV	1647	15	HIV	NEF	216	
RQDILDLWVYHTQGY	1648	15	HIV	NEF	182	
RQEILDLWVYHTQGF	1649	15	HIV	NEF	182	
LSHFLKEKGGLEGLI	1650	15	HIV	NEF	114	
LSFFLKEKGGLDGLI	1651	15	HIV	NEF	114	
LEPWNHPGSQPKTAC		16	HIV	TAT	11	
T	<u>1652</u>	1.5	7713.7	T 4 T	38	
QVCFITKGLGISYGR	<u>1653</u>	15	HIV	TAT		
QLCFLKKGLGISYGR	<u>1654</u>	15	HIV	TAT	38	
PPEESFRFGEEKTTPS	<u> 1655</u>	16	HIV1	gp	81	
CIVYRDGNPYAVCDK	1656	15	HPV	E6	58	
HYCYSLYGTTLEQQY	1657	15	HPV	E6	85	
CYSLYGTTLEQQYNK	1658	15	HPV	E6	87	
NTSLQDIEITCVYCK	1659	15	HPV	E6	22	
VFEFAFKDLFVVYRD	1660	15	HPV	E6	44	
EFAFKDLFVVYRDSI	1661	15	HPV	E6	46	
DLFVVYRDSIPHAAC	1662	15	HPV	E6	51	
FVVYRDSIPHAACHK	1663	15	HPV	E6	53	
NTGLYNLLIRCLRCQ	1664	15	HPV	E6	95	
IRCLRCQKPLNPAEK	1665	15	HPV	E6	103	
PRKLHELSSALEIPY	1666	15	HPV	E6	9	
EIPYDELRLNCVYCK	1667	15	HPV	E6	20	
TEVLDFAFTDLTIVY	1668	15	HPV	E6	40	
VLDFAFTDLTIVYRD	1669	15	HPV	E6	42	
DFAFTDLTIVYRDDT	1670	15	HPV	E6	44	
TIVYRDDTPHGVCTK	1671	15	HPV	E6	51	
WYRYSVYGTTLEKLT	1672	15	HPV	E6	78	
ETTIHNIELQCVECK		15	HPV	E6	20	
SEVYDFAFADLTVVY	1673	15	HPV	E6	40	
VYDFAFADLTVVYRE	1674	15	HPV	E6	42	
DFAFADLTVVYREGN	1675	15	HPV	E6	44	
TVVYREGNPFGICKL	<u>1676</u>	15	HPV	E6	51	
GNPFGICKLCLRFLS	1677	15	HPV	E6	57	
	1678		HPV	E6	80	
NYSVYGNTLEQTVKK	<u> 1679</u>	15				
KKPLNEILIRCIICQ	<u>1680</u>	15	HPV	E6	93	
NEILIRCIICQRPLC	<u> 1681</u>	15	HPV	E6	97	
IRCIICQRPLCPQEK	1682	15	HPV	E6	101	

		HLA-	DR SUPERTY	PE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
CIVYRDCIAYAACHK	1683	15	HPV	E6	53	
NTELYNLLIRCLRCQ	1684	15	HPV	E6	95	
IRCLRCQKPLNPAEK	1685	15	HPV	E6	103	
REVYKFLFTDLRIVY	1686	15	HPV	E6	40	
RIVYRDNNPYGVCIM	1687	15	HPV	E6	51	
NNPYGVCIMCLRFLS	1688	15	HPV	E6	57	
EERVKKPLSEITIRC	1689	15	HPV	E6	89	
IRCIICQTPLCPEEK	1690	15	HPV	E6	101	
EIPLIDLRLSCVYCK	1691	15	HPV	E6	23	
SCVYCKKELTRAEVY	1692	15	HPV	E6	32	
VCLLFYSKVRKYRYY	1693	15	HPV	E6	68	
YYDYSVYGATLESIT	1694	15	HPV	E6	81	
IRCYRCQSPLTPEEK	1695	15	HPV	E6	104	
VYDFVFADLRIVYRD	1696	15	HPV	E6	42	
DFVFADLRIVYRDGN	1697	15	HPV	E6	44	
RIVYRDGNPFAVCKV	1698	15	HPV	E6	51	
GNPFAVCKVCLRLLS	1699	15	HPV	E6	57	
KKCLNEILIRCIICQ	1700	15	HPV	E6	93	
NEILIRCHCORPLC	1700	15	HPV	E6	97	
RTAMFQDPQERPRKL	1701	15	HPV	E6	5	
LFVVYRDSIPHAACH	1702	15	HPV	E6	52	
LTIVYRDDTPHGVCT		15	HPV	E6	50	
LCIVYRDCIAYAACH	1704 1705	15	HPV	E6	52	
YKFLFTDLRIVYRDN		15	HPV	E6	43	
YNFACTELKLVYRDD	1706	15	HPV	E6	46	
LKLVYRDDFPYAVCR	1707	15	HPV	E6	53	
YDFVFADLRIVYRDG	1708 1709	15	HPV	E6	43	
LRIVYRDGNPFAVCK		15	HPV	E6	50	
HEYMLDLQPETTDLY	1710	15	HPV	E7	9	
TLRLCVQSTHVDIRT	1711	15	HPV	E7	64	
IRTLEDLLMGTLGIV	1712	15	HPV	E7	76	
LEDLLMGTLGIVCPI	1713	15	HPV	E7	79	
DLLMGTLGIVCPICS	1714	15	HPV	E7	81	
KATLQDIVLHLEPQN	1715	15	HPV	E7	5	
IDGVNHQHLPARRAE	1716	15	HPV	E7	41	
LRAFQQLFLNTLSFV	1717	15	HPV	E7	83	
FQQLFLNTLSFVCPW	1718	15	HPV	E7	86	
QDYVLDLQPEATDLH	1719	15	HPV	E7	9	
DIRILQELLMGSFGI	1720	15	HPV	E7	75	
IRILQELLMGSFGIV	1721	15	HPV	E7	76	
ELLMGSFGIVCPNCS	1722	15	HPV	E7	81	
KEYVLDLYPEPTDLY	1723	15	HPV	E7	9	
LRTIQQLLMGTVNIV	1724	15	HPV	E7	76	
IQQLLMGTVNIVCPT	1725	15	HPV	E7	79	
QLLMGTVNIVCPTCA	<u>1726</u> 1727	15	HPV	E7	81	
RETLQEIVLHLEPQN		15	HPV	E7	5	
	<u>1728</u>				-	

		HLA-	DR SUPERTY	PE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
LRTLQQLFLSTLSFV	1729	15	HPV	E7	84	
LQQLFLSTLSFVCPW	1730	15	HPV	E7	87	
KDYILDLQPETTDLH	1731	15	HPV	E7	9	
LRTLQQMLLGTLQVV	1732	15	HPV	E7	78	
LQQMLLGTLQVVCPG	1733	15	HPV	E7	81	
QMLLGTLQVVCPGCA		15	HPV	E7	83	
VPTLQDVVLELTPQT	1735	15	HPV	E7	5	
LQDVVLELTPQTEID	1736	15	HPV	E7	8	
QDVVLELTPQTEIDL	1737	15	HPV	E7	9	
CKFVVQLDIQSTKED	1738	15	HPV	E7	68	
VVQLDIQSTKEDLRV	1739	15	HPV	E7	71	
DLRVVQQLLMGALTV		15	HPV	E7	82	
LRVVQQLLMGALTVT	1770	15	HPV	E7	83	
VQQLLMGALTVTCPL	1742	15	HPV	E7	86	
QQLLMGALTVTCPLC	1742	15	HPV	E7	87	
OLLMGALTVTCPLCA		15	HPV	E7	88	
REYILDLHPEPTDLF	1744	15	HPV	E7	9	
TCCYTCGTTVRLCIN	1745	15	HPV	E7	57	
VRTLQQLLMGTCTIV	<u>1746</u>	15	HPV	E7	77	
LOOLLMGTCTIVCPS	1747	15	HPV	E7	80	
MLDLQPETTDLYCYE	1748	15	HPV	E7	12	
VLDLYPEPTDLYCYE	<u>1749</u>	15	HPV	E7	12	
LREYILDLHPEPTDL	<u>1750</u>	15	HPV	E7	8	
HIEFTPTRTDTYACRV	<u>1751</u>	16	Human	B2-µglobulin	67	
LWWVNNESLPVSPRL	<u>1752</u>	15	Human	CEA	177	А
	<u>1753</u>		Human	DRB and	177	^
YEEYVRFDSDVGE	1754	13	Human	CD4 peptide		
EEYVRFDSDVGE		12	Human	DRB and		
A DDDL 10DCDVI EDV	<u>1755</u>	16	Human	CD4 peptide EPO	1	
APPRLICDSRVLERY	<u>1756</u>	15			6	
ICDSRVLERYLLEAK	<u>1757</u>	15	Human	EPO		
VLERYLLEAKEAENI	<u>1758</u>	15	Human	EPO	11	
EHCSLNENITVPDTK	1759	15	Human	EPO	31	
NENITVPDTKVNFYA	<u>1760</u>	15	Human	EPO	36	
VPDTKVNFYAWKRM E	1761	15	Human	EPO	41	
VNFYAWKRMEVGQQ	1701	15	Human	EPO	46	
A	1762					
WKRMEVGQQAVEV WQ	1763	15	Human	EPO	51	
VGQQAVEVWQGLAL	1705	15	Human	EPO	56	
L	1764					
VEVWQGLALLSEAVL	1765	15	Human	EPO	61	
GLALLSEAVLRGQAL	<u>1766</u>	15	Human	EPO	66	
SEAVLRGQALLVNSS	1767	15	Human	EPO	71	
RGQALLVNSSQPWEP	<u>1768</u>	15	Human	EPO	76	
LVNSSQPWEPLQLHV	1769	15	Human	EPO	81	
QPWEPLQLHVDKAVS	1770	15	Human	EPO	86	
LQLHVDKAVSGLRSL	1771	15	Human	EPO	91	

		HLA-	DR SUPERTY	PE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
DKAVSGLRSLTTLLR	1772	15	Human	EPO	96	
GLRSLTTLLRALGAQ	1773	15	Human	EPO	101	
TTLLRALGAQKEAIS	1774	15	Human	EPO	106	
ALGAQKEAISPPDAA	1775	15	Human	EPO	111	
KEAISPPDAASAAPL	1776	15	Human	EPO	116	
PPDAASAAPLRTITA	1777	15	Human	EPO	121	
SAAPLRTITADTFRK	1778	15	Human	EPO	126	
RTITADTFRKLFRVY	1779	15	Human	EPO	131	
DTFRKLFRVYSNFLR	1780	15	Human	EPO	136	
LFRVYSNFLRGKLKL	1781	15	Human	EPO	141	
SNFLRGKLKLYTGEA	1782	15	Human	EPO	146	
KLKLYTGEACRTGDR	1783	15	Human	EPO	152	
APPRLITDSRVLERY	1784	15	Human	EPO	1	Α
ITDSRVLERYLLEAK	1785	15	Human	EPO	6	Α
EHTSLNENITVPDTK	1786	15	Human	EPO	31	Α
KLKLYTGEATRTGDR	1787	15	Human	EPO	152	Α
PQPFRPQQPYPQ	1788	12	Human	gliadin		
PFRPQQPYPQ	1789	10	Human	gliadin		
PQPFRPQQPYP	1790	11	Human	gliadin		
PQPFRPQQP	1791	9	Human	gliadin		
KQPFRPQQPYPQ	1792	12	Human	gliadin		
PKPFRPQQPYPQ	1793	12	Human	gliadin		
PQPFKPQQPYPQ	1794	12	Human	gliadin		
PQPFRKQQPYPQ	1795	12	Human	gliadin		
PQPFRPQKPYPQ	1796	12	Human	gliadin		
PQPFRPQQPKPQ	1797	12	Human	gliadin		
PQPFRPQQPYKQ	1798	12	Human	gliadin		
PQPFRPQQPYPK	1799	12	Human	gliadin		
QFLGQQQPFPPQ	1800	12	Human	gliadin		
FLGQQQPFPPQ	1801	11	Human	gliadin		
LGQQQPFPPQ	1802	10	Human	gliadin		
QFLGQQQPFPP	1803	11	Human	gliadin		
QFLGQQQPF	1804	9	Human	gliadin		
IRNLALQTLPAMCNV	1004	16	Human	gliadin		
Y	1805			-		
NLALQTLPAMCNVY	<u>1806</u>	14	Human	gliadin		
LALQTLPAMCNVY	1807	13	Human	gliadin		
IRNLALQTLPAM	<u>1808</u>	12	Human	gliadin		
IRNLALQTLP	1809	10	Human	gliadin		
EGDAFELTVSCQGGL PK	1810	17	Human	gp100	506	
ESTGMTPEKVPVSEV	1010	18	Human	gp100	370	
MGT	1811					
FPTIPLSRLFDNASL	1812	15	Human	Growth hormone	1	
RLFDNASLRAHRLHQ	1012	15	Human	Growth	8	
	1813			hormone	1.5	
LRAHRLHQLAFDTYQ	1814	15	Human	Growth hormone	15	

		HLA	DR SUPERTYPE			
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
QLAFDTYQEFEEAYI		15	Human	Growth	22	
QEFEEAYIPKEQKYS	<u>1815</u>	15	Human	hormone Growth	29	
QEFERATIFREQUIS	1816	13	riuman	hormone	2)	
IPKEQKYSFLQNPQT		15	Human	Growth	36	
ant oxpossi and	<u>1817</u>	15	Human	hormone Growth	43	
SFLQNPQTSLCFSES	1818	15	Human	hormone	43	
TSLCFSESIPTPSNR		15	Human	Growth	50	
DEETTO OVER IL EL L DI	<u>1819</u>	15	Human	hormone Growth	64	
REETQQKSNLELLRI	1820	13	Human	hormone	04	
SNLELLRISLLLIQS		15	Human	Growth	71	
101 1 1 100W PRIVOR	<u>1821</u>			hormone	70	
ISLLLIQSWLEPVQF	1822	15	Human	Growth hormone	78	
SWLEPVQFLRSVFAN	1022	15	Human	Growth	85	
•	1823			hormone		
FLRSVFANSLVYGAS	1824	15	Human	Growth hormone	92	
NSLVYGASDSNVYDL	1024	15	Human	Growth	99	
	1825			hormone		
SDSNVYDLLKDLEEG	1006	15	Human	Growth	106	
GIQTLMGRLEDGSPR	1826	15	Human	hormone Growth	120	
GIQTEMOREEDOSI R	1827	15	riaman	hormone	120	
RLEDGSPRTGQIFKQ		15	Human	Growth	127	
DTCOIEVOTVEVEDT	<u>1828</u>	15	Human	hormone Growth	134	
RTGQIFKQTYSKFDT	1829	13	riuman	hormone	134	
QTYSKFDTNSHNDDA		15	Human	Growth	141	
	<u>1830</u>			hormone		
TNSHNDDALLKNYGL	1831	15	Human	Growth hormone	148	
ALLKNYGLLYCFRKD	1001	15	Human	Growth	155	
	1832			hormone		
DMDKVETFLRIVQCR	1833	15	Human	Growth hormone	169	
FLRIVQCRSVEGSCGF	1055	16	Human	Growth	176	
-	1834			hormone		
FPTIPLSRLFDNAML	1835	15	Human	Growth	1	Α
RLFDNAMLRAHRLHQ		15	Human	hormone Growth	8	Α
TEL DITTEMENT THE TIME	1836			hormone		
QLAFDTYQEFEQNPQ	1027	15	Human	Growth	22	Α
SFLQNPQTSLCCFRK	<u>1837</u>	15	Human	hormone Growth	43	Α
bi boni qibbeci idi	1838		· · · · · · · · · · · · · · · · · · ·	hormone	-15	••
SNLELLRICLLLIQS	1000	15	Human	Growth	71	Α
ICLLLIQSWLEPVQF	<u>1839</u>	15	Human	hormone Growth	78	Α
ICEEDIQS (FEEL VQI	1840	15	Trumun	hormone	,,,	
NSLVYGASDSNIYDL		15	Human	Growth	99	Α
SDSNIYDLLKDLEEG	1841	15	Human	hormone Growth	106	Α
SUSMITULLADLEEG	1842	13	riuman	hormone	100	Α
DKVETFLRIVQCCGF		15	Human	Growth	169	Α
CELONDOTCI TECES	1843	15	Human	hormone	43	Α
SFLQNPQTSLTFSES	1844	13	riuman	Growth hormone	43	A
TSLTFSESIPTPSNR		15	Human	Growth	50	Α
	1845			hormone		

		HLA-	DR SUPERTY	PE		
	SEQ ID NO.					
Sequence		AA	Organism	Protein	Position	Analog
ALLKNYGLLYTFRKD	1846	15	Human	Growth hormone	155	Α
LLYTFRKDMDKVETF	1040	15	Human	Growth	162	Α
	<u>1847</u>			hormone	160	
DMDKVETFLRIVQTR	1848	15	Human	Growth hormone	169	Α
FLRIVQTRSVEGSTGF	10.10	16	Human	Growth	176	Α
	<u>1849</u>			hormone	40	
HLDMLRHLYQGCQV v	1850	15	Human	Her2/neu	42	
RLRIVRGTQLFEDNYA		17	Human	Her2/neu	98	
L CACCODALACION	<u>1851</u>	1.5	Human	Her2/neu	776	
GVGSPYVSRLLGICL	<u>1852</u>	15		Her2/neu	1166	
TLERPKTLSPGKNGV	<u>1853</u>	15 18	Human Human	Her2/neu Her2/neu	369	
KIFGSLAFLPESFDGDP A	1854	18	Human	Her2/Heu	309	
ELVSEFSRMARDPQ	1855	14	Human	Her2/neu	971	
GEALSTLVLNRLKVG	1856	15	Human	HSP60	280	
AYVLLSEKKISSIQS	1857	15	Human	HSP60	242	
VASLLTTAEVVVTEI	1858	15	Human	HSP60	535	
KCEFQDAYVILLSEKK	1859	16	Human	HSP60	236	
ALSTLVLNRLKVGLQ	1860	15	Human	HSP60	282	
MSYNLLGFLQRSSNC	1861	15	Human	IFN-B	1	
LGFLQRSSNCQCQKL	1862	15	Human	IFN-B	6	
RSSNCQCQKLLWQLN	1863	15	Human	IFN-B	11	
QCQKLLWQLNGRLEY	1864	15	Human	IFN-B	16	
LWQLNGRLEYCLKDR	1865	15	Human	IFN-B	21	
GRLEYCLKDRRNFDI	1866	15	Human	IFN-B	26	
RNFDIPEEIKQLQQF	1867	15	Human	IFN-B	36	
PEEIKQLQQFQKEDA	1868	15	Human	IFN-B	41	
QLQQFQKEDAAVTIY	1869	15	Human	IFN-B	46	
QKEDAAVTIYEMLQN	1870	15	Human	IFN-B	51	
AVTIYEMLQNIFAIF	1871	15	Human	IFN-B	56	
EMLQNIFAIFRQDSS	1872	15	Human	IFN-B	61	
IFAIFRQDSSSTGWN	1873	15	Human	IFN-B	66	
RQDSSSTGWNETIVE	1874	15	Human	IFN-B	71	
STGWNETIVENLLAN	1875	15	Human	IFN-B	76	
ETIVENLLANVYHQR	1876	15	Human	IFN-B	81	
NLLANVYHQRNHLKT	1877	15	Human	IFN-B	86	
VYHORNHLKTVLEEK	1878	15	Human	IFN-B	91	
LEKEDFTRGKRMSSL	1879	15	Human	IFN-B	106	
FTRGKRMSSLHLKRY	1880	15	Human	IFN-B	111	
RMSSLHLKRYYGRIL	1881	15	Human	IFN-B	116	
HLKRYYGRILHYLKA	1882	15	Human	IFN-B	121	
YGRILHYLKAKEDSH	1883	15	Human	IFN-B	126	
HYLKAKEDSHCAWTI	1884	15	Human	IFN-B	131	
KEDSHCAWTIVRVEI	1885	15	Human	IFN-B	136	
CAWTIVRVEILRNFY	1886	15	Human	IFN-B	141	
VRVEILRNFYVINRL	1887	15	Human	IFN-B	146	
RNFYVINRLTGYLRN	1888	15	Human	IFN-B	152	
	1000					

		HLA-	DR SUPERTYPE			
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
MSYNLLGFLQRSSNT	1889	15	Human	IFN-B	1	Α
LGFLQRSSNTQTQKL	1890	15	Human	IFN-B	6	Α
RSSNTQTQKLLWQLN	1891	15	Human	IFN-B	11	Α
QTQKLLWQLNGRLEY		15	Human	IFN-B	16	Α
LWQLNGRLEYTLKDR	1893	15	Human	IFN-B	21	Α
GRLEYTLKDRRNFDI	1894	15	Human	IFN-B	26	Α
HYLKAKEDSHTAWTI	1895	15	Human	IFN-B	131	Α
KEDSHTAWTIVRVEI	1896	15	Human	IFN-B	136	Α
TAWTIVRVEILRNFY	1897	15	Human	IFN-B	141	Α
LGFLQRSSNCQSQKL	1898	15	Human	IFN-B	6	Α
RSSNCQSQKLLWQLN	1899	15	Human	IFN-B	ΙI	Α
QSQKLLWQLNGRLEY	1900	15	Human	IFN-B	16	Α
GIVEQCCTSICSLYQ		15	Human	Insulin alpha	I	
TELESI VOLENIVON	1901	14	Human	chain Insulin al p ha	8	
TSICSLYQLENYCN	1902	14	riuman	chain	0	
GILEQCCTSICSLYQ		15	Human	Insulin alpha	1	Α
CIVEOTTTCITCIVO	<u>1903</u>	15	Human	chain Insulin alpha	I	Α
GIVEQTTTSITSLYQ	1904	13	riuman	chain	1	^
EQTTTSITSLYQLEN	1005	15	Human	Insulin alpha	4	Α
TSICSLYQLENYCG	<u>1905</u>	14	Human	chain Insulin alpha	8	Α
TSICSET QUELTITES	1906		Truman	chain		
TSITSLYQLENYTN	1007	I 4	Human	Insulin alpha	8	Α
TSITSLYQLENYTG	<u>1907</u>	14	Human	chain Insulin alpha	8	Α
7	1908			chain	-	
GIVEQCCCGSHLVEA	1909	15	Human	Insulin alpha-		Α
SLYQLENYCCGERGF	1909	15	Human	beta Insulin alpha-		Α
	<u>1910</u>			beta		
CCTSICSLYQLENYCC	1911	16	Human	Insulin alpha- beta		Α
GSHLVEALYLVCCN	1211	14	Human	Insulin alpha-		Α
	<u>1912</u>			beta		
CCGSHLVEALYLVCC	1913	15	Human	Insulin alpha- beta		Α
FVNQHLCGSHLVEAL	1715	15	Human	Insulin beta	I	
	<u>1914</u>			chain		
QHLCGSHLVEALYLV	1915	15	Human	Insulin beta chain	4	
GSHLVEALYLVCGER		15	Human	Insulin beta	8	
VEALVI VOCEBCEEV	<u>1916</u>	15	Human	chain Insulin beta	12	
VEALYLVCGERGFFY	1917	15	riuman	chain beta	12	
YLVCGERGFFYTPKT		15	Human	Insulin beta	16	
EVNIOUI CCCDI VEAL	<u> 1918</u>	15	Human	chain Insulin beta	I	Α
FVNQHLCGSDLVEAL	1919	13	Tuman	chain		Α.
FVNQHLTGSHLVEAL		15	Human	Insulin beta	I	Α
QHLTGSHLVEALYLV	<u>1920</u>	15	Human	chain Insulin beta	4	A
-	1921			chain		
GSHLVEALYLVTGER	1022	15	Human	Insulin beta	8	Α
VEALYLVCGERGSFY	<u>1922</u>	15	Human	chain Insulin beta	12	Α
	1923			chain		

		HLA-	-DR SUPERTYPE			
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
VEALYLVCGERGFLY		15	Human	Insulin beta	12	A
VEALYLVTGERGFFY	<u>1924</u>	15	Human	chain Insulin beta	12	Α
YLVCGERGFLYTPKT	1925	15	Human	chain Insulin beta	16	Α
ILVCGERGFLIIFRI	1926	13	Hullian	chain	10	^
YLVCGERGFFYTDKT	1927	15	Human	Insulin beta chain	16	Α
YLVCGERGFFYTKPT	1928	15	Human	Insulin beta chain	16	Α
YLVTGERGFFYTPKT	1929	15	Human	Insulin beta chain	16	Α
YLVTGERGFFYTDKT	1930	15	Human	Insulin beta chain	16	Α
YLVTGERGFFYTKPT	1931	15	Human	Insulin beta chain	16	Α
VCGERGFFYTPKTRR	1932	15	Human	Insulin beta chain	18	Α
VTGERGFFYTPKTRR	1933	15	Human	Insulin beta chain	18	Α
MWDLVLSIALSVGCT	1934	15	Human	Kallikrein2	1	
DLVLSIALSVGCTGA	1935	15	Human	Kallikrein2	3	
HPQWVLTAAHCLKK N	1936	15	Human	Kallikrein2	56	
QWVLTAAHCLKKNS O	1937	15	Human	Kallikrein2	58	
GQRVPVSHSFPHPLY	1938	15	Human	Kallikrein2	87	
RVPVSHSFPHPLYNM	1939	15	Human	Kallikrein2	89	
PHPLYNMSLLKHQSL	1940	15	Human	Kallikrein2	97	
HPLYNMSLLKHQSLR	1941	15	Human	Kallikrein2	98	
NMSLLKHQSLRPDED	1942	15	Human	Kallikrein2	102	
SHDLMLLRLSEPAK1	1943	15	Human	Kallikrein2	118	
HDLMLLRLSEPAKIT	1944	15	Human	Kallikrein2	119	
PEEFLRPRSLQCVSL	1945	15	Human	Kallikrein2	162	
PRSLQCVSLHLLSND	1946	15	Human	Kallikrein2	168	
NGVLQGITSWGPEPC	1947	15	Human	Kallikrein2	220	
KPAVYTKVVHYRKWI	1948	15	Human	Kallikrein2	239	
LHLLSNDMCARAYSE	1949	15	Human	Kallikrein2	176	
VGNWQYFFPVIFSKA	1950	15	Human	MAGE3	140	
ESEFQAALSRKVAKL	1951	15	Human	MAGE6	102	
IGHLYIFATCLGLSYD GL	1952	18	Human	MAGE6	172	
VGNWQYFFPVIFSKAS DSLQLVFGIELMEVD		31	Human	MAGE6	140	
PAYEKLSAEQSPPPY	1954	15	Human	MARTI	102	
RNGYRALMDKSLHV	1934	23	Human	MARTI	51	
GTQCALTRR FFKNIVTFFKNIVT	1955 1956	14	Human	МВР		Α
YKSAHKGFKGVDAQ	1750	20	Human	MBP	134	
GTLSKI VDAQGTLSKIFKLGG	<u>1957</u>	20	Human	МВР	144	
RDSRS AC-	<u>1958</u>	23	Human	МВР	1	
ASQKRPSQRHGSKYL ATAST ENPVVHFFKNIVTPR	1959 1960	15	Human	МВР	85	

		HLA-	DR SUPERTY	PE		
	SEQ ID NO.					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
ENPVVAFFKNIVTPR	1961	15	Human	MBP	85	SAAS
ENPVVHAFKNIVTPR	<u>1962</u>	15	Human	MBP	85	SAAS
ENPVVHFFANIVTPR	1963	15	Human	MBP	85	SAAS
ENPVVHFFKNIVTPA	1964	15	Human	MBP	85	SAAS
NPVVHFFKNIVT	1965	12	Human	MBP	86	
HFFKNIVTPRTPPY	1966	14	Human	MBP	90	
NPVVHFFKNIVTPR	1967	14	Human	MBP	86	
LPVPGVLLKEFTVSGN		20	Human	NY-ESO-1	116	
ILTI	<u> 1968</u>	20	Human	NY-ESO-1	161	
WITQCFLPVFLAQPPS GQRR	1969	20	numan	N1-E30-1	101	
DHRQLQLSISSCLQQL		20	Human	NY-ESO-I	141	
SLLM	<u>1970</u>	20	Human	NY-ESO-1	91	
YLAMPFATPMEAELA RRSLA	1971	20	riuman	N I -E3O-1	91	
AAPLLLARAASLSLG	1972	15	Human	PAP	3	
APLLLARAASLSLGF	1973	15	Human	PAP	4	
PLLLARAASLSLGFL	1974	15	Human	PAP	5	
SLSLGFLFLLFFWLD	1975	15	Human	PAP	13	
LLFFWLDRSVLAKEL	1976	15	Human	PAP	· 2I	
DRSVLAKELKFVTLV	1977	15	Human	PAP	27	
AKELKFVTLVFRHGD	1978	15	Human	PAP	32	
RSPIDTFPTDPIKES	1979	15	Human	PAP	47	
FGQLTQLGMEQHYEL	1980	15	Human	PAP	67	
DRTLMSAMTNLAALF	1981	15	Human	PAP	110	
MSAMTNLAALFPPEG	1982	15	Human	PAP	114	
MTNLAALFPPEGVSI	1983	15	Human	PAP	117	
PEGVSIWNPILLWQP	1984	15	Human	PAP	126	
GVSIWNPILLWQPIP		15	Human	PAP	128	
WNPILLWQPIPVHTV	1985	15	Human	PAP	132	
NPILLWQPIPVHTVP	1986	15	Human	PAP	133	
PILLWQPIPVHTVPL	1987	15	Human	PAP	134	
ILLWQPIPVHTVPLS	1988	15	Human	PAP	135	
WQPIPVHTVPLSEDQ	1989	15	Human	PAP	138	
LSGLHGQDLFGIWSK	1990	15	Human	PAP	194	
YDPLYCESVHNFTLP	<u>1991</u>	15	Human	PAP	210	
LPSWATEDTMTKLRE	1992	15	Human	PAP	223	
LRELSELSLLSLYGI	1993	15	Human	PAP	235	
	<u>1994</u>	15	Human	PAP	238	
LSELSLLSLYGIHKQ	<u>1995</u>		Human	PAP	241	
LSLLSLYGIHKQKEK	<u>1996</u>	15	Human			
KSRLQGGVLVNEILN	<u>1997</u>	15		PAP	255	
GGVLVNEILNHMKRA	1998	15	Human	PAP	260	
IPSYKKLIMYSAHDT	<u>1999</u>	15	Human	PAP	277	
YKKLIMYSAHDTTVS	2000	15	Human	PAP	280	
LIMYSAHDTTVSGLQ	2001	15	Human	PAP	283	
DTTVSGLQMALDVYN	2002	15	Human	PAP	290	
ALDVYNGLLPPYASC	2003	15	Human	PAP	299	
LDVYNGLLPPYASCH	2004	15	Human	PAP	300	

		HLA-	DR SUPERTY	PE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
YNGLLPPYASCHLTE	2005	15	Human	PAP	303	
FAELVGPVIPQDWST	2006	15	Human	PAP	356	
TVPLSEDQLLYLPFR	2007	15	Human	PAP	145	
LTELYFEKGEYFVEM	2008	15	Human	PAP	315	
GPVIPQDWSTECMTT	2009	15	Human	PAP	361	
QAHSLERVCHCLGKW	2005	21	Human	PLP	130	
LGHPDK WTTCQSIAFPSKTSASI	2010	20	Human	PLP	181	
QKGRGYRGQHQAHS	2011	20	Human	PLP	121	
LERVCH AATYNFAVLKLMGR GTKF	2012 2013	18	Human	PLP	260	
VATGLCFFGVALFCG	2012	20	Human	PLP	21	
CGHEA	<u>2014</u>	20		DI D	01	
FLYGALLLAEGFYTT GAVRO	2015	20	Human	PLP	81	
SAVPVYIYFNTWTTC QSIAF	2016	20	Human	PLP	171	
TLSVTWIGAAPLILS	2017	15	Human	PSA	5	
SVTWIGAAPLILSRI	2018	15	Human	PSA	7	
VTWIGAAPLILSRIV	2019	15	Human	PSA	8	
SQPWQVLVASRGRAV	2020	15	Human	PSA	31	
GRAVCGGVLVHPQW		15	Human	PSA	42	
V	<u>2021</u>			DC 4	48	
GVLVHPQWVLTAAH C	2022	15	Human	PSA	46	
HPQWVLTAAHCIRNK	2023	15	Human	PSA	52	
QWVLTAAHCIRNKSV	2024	15	Human	PSA	54	
AHCIRNKSVILLGRH	2025	15	Human	PSA	60	
SVILLGRHSLFHPED	2026	15	Human	PSA	67	
VILLGRHSLFHPEDT	2027	15	Human	PSA	68	
GQVFQVSHSFPHPLY	2028	15	Human	PSA	83	
VFQVSHSFPHPLYDM	2029	15	Human	PSA	85	
PHPLYDMSLLKNRFL	2030	15	Human	PSA	93	
SHDLMLLRLSEPAEL	2031	15	Human	PSA	114	
HDLMLLRLSEPAELT	2032	15	Human	PSA	I 15	
TDAVKVMDLPTQEPA	2033	15	Human	PSA	129	
LHVISNDVCAQVHPQ	2034	15	Human	PSA	172	
CAQVHPQKVTKFMLC	2035	15	Human	PSA	180	
GGPLVCNGVLQGITS	2036	15	Human	PSA	210	
GPLVCNGVLQGITSW	2037	15	Human	PSA	211	
NGVLQGITSWGSEPC	2037	15	Human	PSA	216	
RPSLYTKVVHYRKWI	2039	15	Human	PSA	235	
HSLFHPEDTGQVFQV	2040	15	Human	PSA	74	
PRWLCAGALVLAGGF	2040	15	Human	PSM	18	
LGFLFGWFIKSSNEA	2041	15	Human	PSM	35	
LDELKAENIKKFLYN	2042	15	Human	PSM	62	
IKKFLYNFTQIPHLA		15	Human	PSM	70	
KFLYNFTQIPHLAGT	2044 2045	15	Human	PSM	72	
WKEFGLDSVELAHYD	2045	15	Human	PSM	100	
	2040					

		HLA-	DR SUPERTY	PE		
	SEQ					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
LAHYDVLLSYPNKTH	2047	15	Human	PSM	110	
GNEIFNTSLFEPPPP	2048	15	Human	PSM	135	
GKVFRGNKVKNAQL		15	Human	PSM	206	
A GNKVKNAQLAGAKG	2049	15	Human	PSM	211	
V	2050	13	Truman	1 5141	211	
EYAYRRGIAEAVGLP	2051	15	Human	PSM	276	
AEAVGLPSIPVHPIG	2052	15	Human	PSM	284	
AVGLPSIPVHPIGYY	2053	15	Human	PSM	286	
IGYYDAQKLLEKMGG	2054	15	Human	PSM	297	
TGNFSTQKVKMHIHS	2055	15	Human	PSM	334	
TRIYNVIGTLRGAVE	2056	15	Human	PSM	353	
ERGVAYINADSSIEG	2057	15	Human	PSM	444	
GVAYINADSSIEGNY	2058	15	Human	PSM	446	
DSSIEGNYTLRVDCT	2059	15	Human	PSM	453	
NYTLRVDCTPLMYSL	2060	15	Human	PSM	459	
CTPLMYSLVHNLTKE	2061	15	Human	PSM	466	
DFEVFFQRLGIASGR	2062	15	Human	PSM	520	
EVFFQRLGIASGRAR	2063	15	Human	PSM	522	
TNKFSGYPLYHSVYE	2064	15	Human	PSM	543	
YDPMFKYHLTVAQVR		15	Human	PSM	566	
DPMFKYHLTVAQVRG		15	Human	PSM	567	
MFKYHLTVAQVRGG		15	Human	PSM	569	
M	2067			PG1.4		
KYHLTVAQVRGGMV F	2068	15	Human	PSM	571	
VAQVRGGMVFELANS		15	Human	PSM	576	
RGGMVFELANSIVLP	2070	15	Human	PSM	580	
GMVFELANSIVLPFD	2071	15	Human	PSM	582	
VFELANSIVLPFDCR	2072	15	Human	PSM	584	
ADKIYSISMKHPQEM	2073	15	Human	PSM	608	
IYSISMKHPQEMKTY	2074	15	Human	PSM	611	
PQEMKTYSVSFDSLF	2075	15	Human	PSM	619	
TYSVSFDSLFSAVKN	2076	15	Human	PSM	624	
VLRMMNDQLMFLER	2070	15	Human	PSM	660	
A	2077			DG1.4		
LRMMNDQLMFLERA F	2078	15	Human	PSM	661	
RHVIYAPSSHNKYAG	2079	15	Human	PSM	688	
RQIYVAAFTVQAAAE	2080	15	Human	PSM	730	
QIYVAAFTVQAAAET	2081	15	Human	PSM	731	
VAAFTVQAAAETLSE	2082	15	Human	PSM	734	
YISIINEDGNEIFNT	2083	15	Human	PSM	127	
ISIINEDGNEIFNTS	2084	15	Human	PSM	128	
EDFFKLERDMKINCS	2085	15	Human	PSM	183	
FFKLERDMKINCSGK	2085	15	Human	PSM	185	
GVILYSDPADYFAPG	2087	15	Human	PSM	224	
GAAVVHEIVRSFGTL	2088	15	Human	PSM	391	
NSRLLQERGVAYINA	2089	15	Human	PSM	438	
	2002					

		HLA-	-DR SUPERTYPI	3		
	SEQ ID NO.					
Sequence	ID NO.	AA	Organism	Protein	Position	Analog
VAYINADSSIEGNYT	2090	15	Human	PSM	447	
DQLMFLERAFIDPLG	2091	15	Human	PSM	666	
KSNFLNCYVSGFHPSD		16	Human B2-		19	
AC-	<u>2092</u>	18	μglobulin IEd	MHC derived	Unknown	
NPDAENWNSQFEILE		10	TIAL .	wiric derived	Cikilowii	
DAA	2093				0.5	
EYLILSARDVLAVVS	2094	15	M. leprae		85	
YKTIAYDEEARR	2095	12	MT	110000	3	
GEALSTLVVNKIRGT	2096	15	Mycobacteria	HSP60	254	
PYILLVSSKVSTVKD	2097	15	Mycobacteria	HSP60	216	
EAVLEDPYILLVSSK	2098	15	Mycobacteria	HSP60	210	
IAGLFLTTEAVVADK	2099	15	Mycobacteria	HSP60	507	
ALSTLVVNKIRGTFK	2100	15	Mycobacteria	HSP60	256	
MKHILYISFYFILVN	2101	15	Pf	LSA1	1	
KSLLSTNLPYGRTNL	2102		Pf	SSP2	116	
HFFLFLLYILFLVKM	2103	15	Pf		13	
LFLLYILFLVKMNAL	2104	15	Pf		16	
ILFLVKMNALRRLPV	2105	15	Pf		21	
MNALRRLPV1CSFLV	2106	15	Pf		27	
SAFLESQSMNKIGDD	2107	15	Pf		79	
LKELIKVGLPSFENL	2108	15	Pf		132	
FENLVAENVKPPKVD	2109	15	Pf		143	
PATYGIIVPVLTSLF	2110	15	Pf		158	
YGIIVPVLTSLFNKV	2111	15	Pf		161	
LLKIWKNYMKIMNHL	2112	15	Pf		28	
MTLYQIQVMKRNQK		15	Pf		43	
Q	<u>2113</u>	15	Pf		57	
QKQVQMMIMIKFMG V	2114	13	FI		37	
MIMIKFMGVIYIMII	2115	15	Pf		63	
GVIYIMIISKKMMRK	2116	15	Pf		70	
LYYLFNQHIKKELYH	2117	15	Pf		285	
HFNMLKNKMQSSFFM		15	Pf		299	
LDIYQKLYIKQEEQK	2119	15	Pf		353	
QKKY1YNLIMNTQNK	2120	15	Pf		366	
YEALIKLLPFSKRIR	2121	15	Pf		381	
ENEYATGAVRPFQAA	2122	15	Pf		2	
NYELSKKAVIFTPIY	2123	15	Pf		27	
QKILIKIPVTKNIIT	2124	15	Pf		108	
KCLVISQVSNSDSYK	2125	15	Pf		156	
SKIMKLPKLPISNGK	2126	15	Pf		202	
FIHFFTWGTMFVPKY	2127	15	Pf		220	
LCNFKKNIIALLIIP	2128	15	Pf		242	
KKNIIALLIIPPKIH	2129	15	Pf		246	
ALLIIPPKIHISIEL	2130	15	Pf		251	
SMEYKKDFLITARKP	2131	15	Pf		274	
KSKFNILSSPLFNNF	2132	15	Pf		7	
	2172					

		HLA-	DR SUPERTYP	E		
	SEQ ID NO.					
Sequence	ID NO.	AÁ	Organism	Protein	Position	Analog
FKKLKNHVLFLQMM		15	Pf		173	
N KNHVLFLQMMNVNL	<u>2133</u>	15	Pf		177	
Q VLFLQMMNVNLQKQ	2134	15	Pf		180	
L	2135					
NVNLQKQLLTNHLIN	2136	15	Pf		187	
QKQLLTNHLINTPKI	2137	15	Pf		191	
NHLINTPKIMPHHII	2138	15	Pf		197	
YILLKKILSSRFNQM	2139	15	Pf		239	
FNQMIFVSSIFISFY	2140	15	Pf		250	
KVSCKGSGYTFTAYQ		17	Rheumatiod	Variable		
MH	<u>2141</u>	20	vector	region	1	
IAKVPPGPNITAEYGD KWLD	2142	20	Rye grass	Lolp1	'	
TAEYGDKWLDAKST		20	Rye grass	Lolp1	11	
WYGKPT	<u>2143</u>	20	D	T -1-1	21	
AKSTWYGKPTGAGPK DNGGA	2144	20	Rye grass	Lolpi	21	
GAGPKDNGGACGYK		20	Rye grass	Lolp1	31	
DVDKAP	2145	•			51	
FNGMTGCGNTPIFKD GRGCG	2146	20	Rye grass	Lolp1	31	
PIFKDGRGCGSCFEIK	21.10	20	Rye grass	Lolp1	61	
CTKP	2147		_ 1			
SCFEIKCTKPESCSGE AVTV	2148	20	Rye grass	Lolp1		
AFGSMAKKGEEQNVR		20	Rye grass	Lolp1	111	
SAGEL	2149			Ī		
TPDKLTGPFTVRYTTE	2150	20	Rye grass	Lolp1	201	
GGTK VRYTTEGGTKSEVED	2130	20	Rye grass	Lolp1	211	
VIPEG	2151			·		
TCVLGKLSQELHKLQ	2152	15	Salmon	Calcitonin	6	
KLSQELHKLQTYPRT	2153	15	Salmon	Calcitonin	11	
LHKLQTYPRTNTGSG	2154	15	Salmon	Calcitonin	16	
KLQTYPRTNTGSGTP	2155	15	Salmon	Calcitonin	18	
CCVLGKLSQELHKLQ	2156	15	Salmon	Calcitonin	7	Α
CSNLSTCVLGKLSQE	2157	15	Salmon	Calcitonin	1	Α
TSNLSTTVLGKLSQE	2158	15	Salmon	Calcitonin	1	Α
TTVLGKLSQELHKLQ	2159	15	Salmon	Calcitonin	6	Α
DIAAKYKELGY	2160	11	Sperm whale	Myoglobin	141	
ALVRQGLAKVA	.2161	11	Staph.	Nase	102	
PATLIKAIDGDTVKLM		20	Staph.	Nase	11	
YKGQ	2162		-			
TPETKHPKKGVEKYG	2162	20	Staph.	Nase	41	
PEASA VEKYGPEASAFTKKM	<u>2163</u>	20	Staph.	Nase	51	
VENAK	2164		•			
FTKKMVENAKKIEVE	2165	20	Staph.	Nase	61	
FDKGQ YIYADGKMVNEALVR	<u>2165</u>	20	Staph.	Nase	91	
QGLAK	2166	20	олари.	.1430	71	
HEQHLRKSEAQAKKE		20	Staph.	Nase	121	
KLNIW QAKKEKLNIWSEDNA	2167	19	Staph.	Nase	131	
DSGQ	2168	17	o.apii.	.1430		
•						

	CEO	nLA-	DR SUPERTY	T.E.		
	SEQ ID NO.		0	Protein	Position	4
Sequence YFNNFTVSFWLRVPK	10160	15	Organism TetTox	rrotein	947	Analog
FSYFPSI	2169	7	TetTox		593	Α
YSFFPSI	2170	7	TetTox		593	A
YSYFPSIR	2171	8	TetTox		593	A
DPNANPNVDPNANPN	2172	117	Unknown	(MAP)=(T1B	373	Α.
VNANPNANPNANP(X		117	Clikilowii)4		
4)	2173			· .		
QKWAAVVVPS	2174	10	Unknown	ClassI A2	242	
TWQLNGEELIQDMEL VETRPAG	2175	22	Unknown	ClassI Kb 216-237	216	
PEFLEQRRAAVDTYC	2176	15	Unknown	IEBs2		
STORKUSP33	2170		Unknown	RAGE		
DYSYLODSDPDSFQD	2178	15	Unknown	Tyrosinase	448	
DFSYLQDSDPDSFQD	2179	15	Unknown	Tyrosinase	448	SAAS
QNILFSNAPLGPQFP	2180	15	Unknown	Tyrosinase	56	SAAS
QNILLSNAPLVPQFP	2181	15	Unknown	Tyrosinase	56	SAAS
DYSYLQDSDPDSFQD	2182	15	Unknown	Tyrosinase	448	
KYVKQNTLKLAT	2183	11	unknown			
P(X)KQNTLKLAT	2184	13	unknown			Α
EEDIEIIPIQEEEY	2185	14		CD20	249	Α
HQAISPRTLNSPAIF	2186	15				
YTDVFSLDPTFTIETT	2187	16				
YAGIRRDGLLLRLVD	2188	15				Α
LFFYRKSVWSKLQS1	2189	15				
RPIVNMDYVVGARTF	2102	20				
RREKR	2190					
RPGLLGASVLGLDDI	2191	15				
LYFVKVDVTGAYDTI	2192	15				
FAGIRRDGLLLRLVD	2193	15				
AKTFLRTLVRGVPEY	2194	15				
YGAVVNLRKTVVNFP	2195	15				
GTAFVQMPAHGLFPW	<u>2196</u>	15				
WAGLLLDTRTLEVQS	2197	15				
RTSIRASLTFNRGFK	<u>2198</u>	15				
RVIKNSIRLTL	2199	11				
PVIKNSIKLRL	2200	11				
ATSTKKLHKEPATLIK AIDG	2201	21				

TABLE 26

	HLA-DR SUPERTYPE										
Sequence	SEO ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB:		
AC-					>900000	500000		25000			
NPTKHKWEAAHV AEOLAA	1543										
DDŶVKQYTKQYTK		50000		160	500000		12500				
QNTLKK AAAKAAAAAAYA	<u>1544</u>	833	>900000	229	500000		12500				
A	1545										
AC- AAAKAAAAAAYA		625		348							
A	<u>1546</u>										
(20)AYA(20)A(20)A(20)K(20)A(20)	<u>1547</u>	50000		250	500000		8333				
AC-	1011	50000		381							
AAAKATAAAAYA A	1548										
AC-	1040	50000		421							
AAAKAAAAAAFA A	1549										
AC-	1373	5000		444	500000						
AAAKATAAAA(10) AA	<u>1550</u>										
AC-	1550	1250		286	25000						
AAAKATAAAA(23)	4554										
AA AAKAAAAAAA(10)	<u>1551</u>	2500		>888.89							
AA	<u>1552</u>	2.0		0.54	2770						
AAYAAAATAKAA A	1553	. 3.9		0.54	2778						
AALAAAAAKAA		1.9		12	152		1316				
A AAEAAAATAKAAA	<u>1554</u>	2500		667	500000						
AAYJJAAAAKAAA	1555	50000		533	500000						
AAYAAAAJJKAAA	<u>1556</u>	1250		308	500000						
AFLRAAAAAAFAA	<u>1557</u>	50000		400	500000						
AFLRQAAAAAFAA	<u>1558</u>	2500		1000	25000						
Y	<u>1559</u>		1063	0.19	6.2		67				
AAFAAAKTAAAFA	<u>1560</u>	1.3	1003				34				
YAAFAAAKTAAAF A	1561	0.74		0.13	5.0		34				
AALKATAAAAAA	1562	50000		800	500000						
YAR(15)ASQTTLKA		1.5		0.46	5.2		1196				
KT YARF(33)QTTLKAK	<u>1563</u>	50000		889	16667						
T PKYFKQRILKFAT	1564	1667		400	1042						
PKYFKQGFLKGAT	<u>1565</u>	50000		800	500000						
PKYGKQIDLKGAT	<u>1566</u>	50000		444	500000						
AAFFFFFGGGGGA	<u>1567</u>	50000		800	500000						
AADFFFFFFFDA	<u>1568</u>	1250		286	500000						
AAKGIKIGFGIFA	<u>1569</u>	50000		471	500000						
AAFIFIGGGKIKA	<u>1570</u>	50000		195	500000						
	<u>1571</u>			200	25000						
AAKIFIGFFIDGA	<u>1572</u>	1250									
AAFIGFGKIKFIA	<u>1573</u>	50000		242	500000						
AAKIGFGIKIGFA	1574	50000		889	500000						
AAFKIGKFGIFFA	<u>1575</u>	50000		615	500000						
AADDDDDDDDDD A	1576	50000		667	500000						
(43)AAIGFFFFKKGI		50000		258	500000						
A	<u>1577</u>										

Sequence				н	A-DR SUP	RTYPE			
G3)AADFGIFKIGKF 50000 381 500000	Sequence	SEQ ID							
(43)AADFGIFIDFIIA (43)AAFIGFIFIKD (43)AAFIGFIFIKD (43)AAFIGFIGKIGFI (43)AAFIGFIKIGFI (43)AAFIGFIKIGFI (54)AAFIGFIKIGFI (54)AAAAAAAAAA (55)AAAAAAAAAAA (55)AAAAAAAAAA									
(43)AARGGFFRKD 1580 50000 1000 500000	A (43)AADFGIFIDFIIA		50000		235	500000			
(43)AAKIGFGKIKFI (43)AAKIGFGKIGF (43)AAKIGFGKIGF (43)AAKAAAAAAA 550000 1000 500000 276 500000 1582 500000 276 500000 727 5000000 AAAKAAAAAAAAA 5584 50000 727 500000 727 500000 AAAKAAAAAAAAA 5885 50000 50000 500000 500000 500000 500000 500000 500000 500000 600000 600000 600000 6000000 6000000	(43)AAIGGIFIFKKD		50000		800	500000			
(43)AAKIGFGIKIGF (43)AAKIGFGIKIAG (43)AAKIGFGIKAAA (44)AAAIGAAAAAAA (45)AAKIGFGIKAAAA (46)AAKIAAAAAAA (46)AAKIAAAA (46)AAKIAAAAA (46)AAKIAAAA (46)AAKIAAAAAAA (46)AAKIAAAAAAAAA (46)AAKIAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	A (43)AAFIGFGKIKFI		50000		1000	500000			
(43)AAFKICKFGIFF AAAKAAAAAAA 1588 AAAKAAAAAAAA 1588 AAAKAAAAAAAA 1588 AAAAAAAAAAA 1588 AAAAAAAAAA	(43)AAKIGFGIKIGF		50000		1000	500000			
AAAKAAAAAAA 1584 50000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 500000 727 72846 728	(43)AAFKIGKFGIFF		50000		276	500000			
AAAKAAAAAAFA	AAAKAAAAAAA		>1666.67		>347.83	12500			
AAAKAAAAAFAA	AAAKAAAAAAAF		50000		727	500000			
AAAKAAAAAAAAA	AAAKAAAAAAFA		50000		235	25000			
FAAAAAAAAAAA	AAAKAAAAFAAA		50000		533	500000			
AAAAAAAAAAA	FAAAAAAAAAA		1667		200	8333			
AAAAAAAAAAAA 1590 50000 1000 500000	AAAAAAAAAAA		50000		500	500000			
AAANAAAAAAA 591 50000 615 500000 AAAAAAAAAAAA 1591 50000 533 500000 AAAAAAAAAAAA 1592 50000 3235 500000 AAAAAAAAAAAA 1593 50000 364 500000 AFAAAKTAA 1595 50000 571 500000 AFAAAKTAA 1595 5000 371 500000 AFAAAKTAA 1596 5000 571 500000 AFAAAKTAA 1599 1.6 0.35 3.8 8333 AAAAAAAAAAA 1599 4.2 0.31 4.3 250000 AAAAAAAAAAAA 1600 455 1.3 37 250000 AAAAAAAAAAAA 1600 5000 571 1852 AAAAAAAAAAAA 1601 5000 571 1852 AAAAAAAAAAAAA 1602 3333 727 500000 AAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	AAAAAAAAAAN		50000		1000	500000			
AAAAAAAAAAA A 59000 533 500000 AAAAAAAAAAAA A 1592 50000 364 500000 AFAAAAAAAAAA A 1595 50000 571 500000 AFAAAKTAA 1596 24 0.78 5.2 1786 VAR(15A)RQTTLKAA A 1599 4.2 0.31 4.3 250000 (16A)RQTTLKAAAA 1599 4.2 0.31 4.3 250000 (16A)RQTTLKAAAA 1600 571 1852 (39)AAAATKAAAA 1602 3333 727 500000 (39)AAAATKAAAA 1602 3333 727 500000 (35)AAAATKAAAA 1602 3333 727 500000 (35)AAAATKAAAA 1602 3333 727 500000 (35)AAAATKAAAA 1602 399 0.45 54 96 AA(14)AAAKTAAA 1600 590 667 5556 AA(14)AAAKTAAAA 1600 590 0.45 54 96 AA(14)AAAKTAAA 1600 5000 667 550000 AA(14)AAAKTAAA 1600 5000 667 500000 AA(14)AAAAKTAAAA 1600 5000 667 500000 AA(14)AAAAAAAA 1600 5000 667 500000 AA(14)AAAAAAAAAA 1600 5000 667 500000 AA(14)AAAAAAAAAAA 1600 5000 667 500000 AA(14)AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	AAANAAAAAAA		50000		615	500000			
AAAAAAAAAAA 1593 50000 235 500000 ASAAAAAAAAAA 1594 500000 364 500000 AFAAAKTAA 1595 50000 571 500000 ARACIAS)SQTTLKA KT YAR(ISA)RQTTLKA AA (ISA)RQTTLKAA AI (ISA)RQTTLKAAA 1599 4.2 0.31 4.3 250000 (IGA)RQTTLKAAAA 1600 455 1.3 37 250000 (IGA)RQTTLKAAA 1601 3333 727 500000 (SS)AAAATKAAAA 1602 3333 727 500000 (SS)AAAATKAAAA 1602 3333 727 500000 (SS)AAAATKAAAA 1602 3000 242 2632 (SS)AAAATKAAAA 1603 2000 242 2632 (SS)AAAATKAAAA 1604 2500 667 5556 AA(I4)AAAKTAAAA 1605 39 0.45 54 96 AA(I4)AA(SS)ATKAA AA AA(I4)AA(SS)ATKAA AA AA(I4)AA(SS)ATKAA AA(I4)AA(SS)ATKAA AA(I4)AA(SS)ATKAAA AA(I4)AA(SS)ATKAAAA AA(I4)AA(SS)ATKAAAAA AA(I4)AA(SS)ATKAAAAA AA(I4)AA(SS)ATKAAAAA AA(I4)AA(SS)ATKAAAAAA AA(I4)AA(SS)ATKAAAAA AA(I4)AA(SS)ATKAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	AAAAAAAAAAA		50000		533	500000			
ASAAAAAAAA A 1595 50000 364 500000 AFAAAKTAA 1595 50000 571 500000 AFAACKTAA 1596 2.4 0.78 5.2 1786 XT YAR(15A)RQTTLKA AA (15A)RQTTLKAA AB (16A)RQTTLKAAA AA (16A)RQTTLKAAA AB (16A)RATAAAAAA AB (16A)RATKAAAA AB (16B) 3333 727 500000 AB (3S)AAAATKAAAA AB (16A)RATKAAAA AB (16B) 3000 571 1852 AB (3B)AAATKAAAA AB (16B) 3000 571 1852 AB (3B)AAATKAAAA AB (16A)RATKAAAA AB (16B) 3000 667 5556 AB (16A)RATKAAAA AB (16A)RATKAAAA AB (16A)RATKAAAA AB (16B) 3000 667 5556 AB (16A)RATKAAAA AB AB (16A)RATKAAAA AB	AAAAASAAAAAA		50000		235	500000			
AFAAAKTAAA AFAAAKTAAA AFAAAKTAAA AFAAAKTAAAA AFAAAKTAAAA AA(14)AA(35)AKAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAAKTAAAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAA AA(14)AAA AA(14)AAAKTAAAA AA(14)AAA AA(14)AAAAAAAAAAAA AA(14)AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	ASAAAAAAAAA		50000		364	500000			
YARFICALITICRAR YAR(ISA)SQITLKA KT YAR(ISA)RQITLKA TARRISA)RQITLKAAA 1596 160 1598 160 1598 160 160)RQITLKAAA 2.4 1598 160 1598 160 1598 160 160)RQITLKAAA 1598 160 1598 160 1599 160)RQITLKAAA 1598 160 1598 160 160 160 160 160 160 160 160 160 160			50000		571	500000			
YAR(ISA)SQITILKA KI 1597 YAR(ISA)RQITILKA AI 1598 (ISA)RQITILKAAA 1599 (ISA)RQITILKAAA 1599 (ISA)RQITILKAAA 1599 (ISA)RQITILKAAA 1599 (ISA)RQITILKAAA 1599 (ISA)RQITILKAAA 1599 (ISA)RATKAAAAA 1500 (ISA)RATKAAAA 1500 (ISA)RAATKAAAA 1500 (ISA)RATKAAAAA 1500 (ISA)RATKAAAAA 1500 (ISA)RAATKAAAAA 1500 (ISA)RATKAAAAA 1500 (ISA)RATKAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA	YARFLALTTLRAR		0.98		0.28	3.4			
YAR(IAS)RQTTLKA AA (ISA)RQTTLKAAA 1599 (ISA)RQTTLKAAA 1599 (ISA)RQTTLKAAA 1599 (ISA)RQTTLKAAA 1599 44.2 1590 45.5 1.3 37 250000 (ISA)AAATKAAAA 1601 33333 727 50000 371 1852 (S2)AAAATKAAAA 1601 2000 242 2632 (S2)AAAATKAAAA 1603 2000 242 2632 (S2)AAAATKAAAA 1603 39 0.45 54 96 4(14)AAAKTAAAA 1605 39 0.45 54 96 4(14)AAAKTAAAA 1606 39 0.45 54 96 4(14)AAAKTAAAA 1607 1608 39 0.45 54 96 4(14)AAAKTAAAA 1608 1609 1609 1609 1609 1609 1609 1609 1609	YAR(15A)SQTTLKA		2.4		0.78	5.2		1786	
(16A)RQTTLKAAA	YAR(15A)RQTTLKA		1.6		0.35	3.8		8333	
(16A)RQTTLKAAA 1500 455 1.3 37 250000 (46)AAKTAAAFA 1501 5000 571 1852 (39)AAAATKAAAA 1502 3333 727 500000 (52)AAAATKAAAA 1603 2000 242 2632 (52)AAAATKAAAA 1604 2500 667 5556 A(14)AAAKTAAA 1605 39 0.45 54 96 A(14)AAAKTAAA 1605 39 0.45 50000 500000 AA 1606 5000 667 25000 500000 500000 49AAAKTA(A(72) 1608 5000 533 500000 500000 667 500000 499AAAKTA(49)AA 1610 5000 533 500000 667 500000 667 500000 499AAAKTA(49)AA 1610 5000 533 500000 667 500000 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 <t< td=""><td></td><td></td><td>4.2</td><td></td><td>0.31</td><td>4.3</td><td></td><td>250000</td><td></td></t<>			4.2		0.31	4.3		250000	
(46)AAKTAAAFA (39)AAATKAAAA (3902) (35)AAATKAAAA (3602) (35)AAATKAAAA (3602) (35)AAATKAAAA (3602) (35)AAATKAAAA (3602) (35)AAATKAAAA (3603) (3604) (3	(16A)RQTTLKAAA		455		1.3	37		250000	
(39)AAATKAAAA	(46)AAKTAAAFA		5000		571	1852			
(\$3)AAATKAAAA	(39)AAAATKAAA		3333		727	500000			
(\$5)AAATKAAAA	(52)AAAATKAAAA		2000		242	2632			
A(14)AAAKTAAA 1605 39 0.45 54 96 A(14)A(3(3)ATKAA A(14)A(3(3)ATKAA A(14)A(3(3)ATKAA A(14)A(3(3)ATKAA A(14)A(3(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3)TKAA A(15)A(3(3)TKAA A(15)A(3)TKAA A(15)A(3)TKAA A(15)A(3)TKAA A(15)A(15)A(15)A(15)A A(15)A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A A(15)A(15)A(15)A A(15)A(15)A A(1	(55)AAAATKAAAA		2500		667	5556			
AA(14)A(35)ATKAA AA AA(14)AA(36)TKAA AA AA(14)AA(36)TKAA AA AA(14)AA(36)TKAA AA AA AFAAAKTAA(72) 1608 5000 667 25000 49)AAAKT(64)AA 1609 50000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 667 500000 533 500000 667 500000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 50000 500000 500000 500000 500000 500000 5000000			39		0.45	54		96	
AA(14)AA(36)TKAA AA	AA(14)A(35)ATKAA								
AFAAKTAA(72) 1808 5000 533 500000 (49)AAAKT(64)AA 1610 50000 667 500000 HQAISPRITNGFGP GSPAIF YAAFAAKTAAF TEGRCLHYTVDKS KFK AWYAWNNKCK 1613 50000 667 500000 1610 50000 533 500000 1611 1.9 0.82 7.0 1612 1667 200 500000 >2250000 KFK AWYAWNNKCK 1613 50000 667 500000 >212500 18333 \$8333.33	AA(14)AA(36)TKAA		50000		667	25000			
(49)AAAKT(64)AAA 1609 50000 667 500000 (49)AAAKT(64)AA 1610 50000 533 500000 HQAISPRTINGPOP GSPAIF 1611 1555 728464 12089 2056 3107 5081 YAAFAAAKTAAAF 1612 1.9 0.82 7.0 TEGRCLHYTVDKS KPK 1613 200 500000 >250000 KPX 1613 50000 667 500000 >12500 IVSDONGMNAWV 1250 18371 1000 8333 >8333.33			5000		533	500000			
(49)AAKT7(64)AA 1610 50000 533 500000 HQAISPRILNGPGP GSPAIF YAAFAAKTAAFA TEGRCLHYTVDKS KRX AWYAWRNRCK 1612 163 1657 28464 12089 2056 3107 5081 1612 KRX AWYAWRNRCK 1612 1613 169 282 7.0 200 2500000 2500000 2500000 250000	(49)AAAKT(64)AAA		50000		667	500000			
HQAISPRTLNGPGP GSPAIF 1555 728464 12089 2056 3107 5081	(49)AAAKTA(64)AA		50000		533	500000			
GSPAIF VAAFAAAKTAAAF A TEGRCLHYTVDKS KPK AWYAWRNRCK 1612 1667 200 200 500000 500000 >250000 AWYAWRNCK WSDONGMNAWY 1613 1614 50000 1617 667 1619 500000 167 >12500 500000 WSDONGMNAWY 1250 18371 118371 1000 118331 1000 8333 18333.33 >8333.33		1610	1555	728464	12089	2056	3107	5081	
A 612 500000 5250000 5250000 5250000 5250000 5250000 5250000 5250000 5250000 5250000 5250000 52500000 52500000 525000000 5250000000000	GSPAIF	<u>1611</u>							
AWVAWRNRCK 1614 50000 667 500000 >12500 1VSDGNGMNAWV 1250 18371 1000 8333 >8333.33	A TEGRCLHYTVDKS		1667		200	500000		>250000	
IVSDGNGMNAWV 1250 18371 1000 8333 >8333.33			50000		667	500000		>12500	
			1250	18371	1000	8333		>8333.33	

HLA-DR SUPERTYPE									
Sequence	SEQ ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB *090
PHHTALRQAILSW		1250		166	1773		14434		
GELMTLA WMYYHGQRHSDE HHH	<u>1616</u> 1617	50000	>900000	727	500000		>250000		
YIVMSDWTGGA		50000	13416	222	500000		12500		
АНААНААНААНА АНАА	1618 1619	263		80000	500000		>250000		
MDIDPYKEFGATV	1015	1563		170			6609		
ELLSFLPSDFFP GMLPVCPLIPGSST	<u>1620</u>	1250	>900000	400	1220		250000		
ISTGP LGFFPDHQLDPAFR	1621	1667	12027	333	2941		250000		
ANT GYKVLVLNPSV	1622	16	72407	27	2116	145	1516	115	8789
LMAFTAAVTS	<u>1623</u>	2511	>73952.34	321	20577	627	240	>40562.91	160
FALWRVSAEEY	1624	>5279.83	88348	342	569	72	927	1433	517
	<u>1625</u>								
ALWRVSAEEY	1626	>6337.14	>76595.74	6543	6669	>35315.99	7954	4099	698
EEYVEIRQVGDFH	1627	>1957.71	74884	>5365.53	11627	26	11323	13890	1115
VGGVYLLPRRGPR LGV	1628	177	236639	22323	12756	2764	351		
VGGAYLLPRRGPR LGV	1629	131	308534	26164	125056	>12230.45	703		
/GGVALLPRRGPR _GV	1630	849	326288	48233	23669	>12230.45	61558		
VGGVYALPRRGPR LGV	<u>1631</u>	134	348950	25750	30504	>12230.45	749		
VGGVYLAPRRGPR LGV	1632	746	202660	33672	>116550.12	>12230.45	878		
VGGVYLLARRGPR LGV	1633	60	23276	485	4396	2199	595		
VGGVYLLPARGPR LGV	1634	12	68070	3644	3213	4579	49		
VGGVYLLRRAGPR LGV	<u>1635</u>	202	39751	12252	32330	6432	433		
GAPLGGAARALAH GV	1636	690	3145	10408	19762	>13044.97	10773		
GAALGGAARALAH GV	<u>1637</u>	1081	26944	21362	60600	>13044.97	29786		
GAPLAGAARALAH GV	<u>1638</u>	588	2983	39885	19692	>13044.97	8178		
GAPLGAAARALAH GV	1639	226	17703	10255	52041	>13044.97	6490		
GAPLGGLARALAH GV	1640	537	351525	13941	6564	>13044.97	66		
GAPLGGALRALAH GV	<u>1641</u>	68	>486486.49 82088	14977	977 1272	1271 >3365.21	1418 31907		
GAPLGGAAAALAH GV	1642	147 398	22959	5472 14984	21017	>3365.21	57549		
GAPLGGAARLLAH GV GAPLGGAARAAAH	<u>1643</u>	797	377964	25279	>110132.16	>3365.21	31308		
OV OAPLGGAARALAA	<u>1644</u>	541	23298	11270	16747	>3365.21	7419		
GV FPDWQNYTPGPGT	1645	13766	>223880.6	23394	>109170.31	>10101.01	59625	592	3013
RF RFPLTFGWCFKLVP	<u>1646</u>	5913	406579	316	21384	121	4100	748	1848
V RQDILDLWVYHTQ	<u>1647</u>	2390	98327	1202	1624	1136	1628	5039	1665
SY RQEILDLWVYHTQ	1648	1050	10530	5928	1414	3362	3052	2730	3679
GF LSHFLKEKGGLEGL	<u>1649</u>	537	>340909.09	2442	86814	2114	13676	1561	2319
LSFFLKEKGGLDGL	<u>1650</u>	172	>340909.09	1275	>109170.31	983	19957	1127	3501
	1651								3446

			HL.	A-DR SUPI	ERTYPE				
	SEQ ID	DRBI	DRB1	DRB1 *0401	DRBI	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
Sequence OVCFITKGLGISYG	NO.	*0101	*0301 166744	1529	*0404 1391	295	91	41	296
Ř QLCFLKKGLGISYG	<u>1653</u>	185	158381	4436	1613	443	3634	40	200
R PPEESFRFGEEKTTP	1654	>2500	>900000	267	500000		>12500		
S CIVYRDGNPYAVC	<u>1655</u>	8464		147	1084	3473	>17182.13		31865
DK HYCYSLYGTTLEQ	<u>1656</u>	546		1127	9713	76	9858		12359
QY CYSLYGTTLEQQY	1657	1086		1317	2836	71	>9964.13		25989
NK NTSLQDIEITCVYC	<u>1658</u>	>12106.54		10930	6143	4584	>17182.13		30884
K VFEFAFKDLFVVYR	<u>1659</u>	6716		1059	2156	120	11583		16797
D EFAFKDLFVVYRDS	<u>1660</u>	8944		2220	11721	33	3688		1882
I DLFVVYRDSIPHAA	<u>1661</u>	1186		82	218	3591	5213		2374
C FVVYRDSIPHAACH	<u>1662</u>	587	200	10	87	704	5085		2122
K NTGLYNLLIRCLRC	1663	127	13429	686	358	258	6743		4759
Q IRCLRCQKPLNPAE	<u>1664</u>	7240		6334	8464	1229	16787		32024
K PRKLHELSSALEIPY	1665	156	16146	5276	694	80	103		213
EIPYDELRLNCVYC	1666	3299		15532	11292	7321	>35612.54		>39432.18
K TEVLDFAFTDLTIV	1667	2073	1542	185	1083	871	1432		349
Y VLDFAFTDLTIVYR	<u>1668</u>	354	30	313	6061	721	230		252
D DFAFTDLTIVYRDD	1669	463	23	80	3373	40	725		1443
T TIVYRDDTPHGVCT	<u>1670</u>	3798		22	1269	>9753.59	>35612.54		>39144.05
K WYRYSVYGTTLEK	<u>1671</u>	163	26561	249	3448	8.5	107		284
LT ETTIHNIELQCVEC	1672	3623		1996	3327	6561	>35612.54		>39432.18
K SEVYDFAFADLTV	<u>1673</u>	31	2996	260	2180	101	1850		174
VY VYDFAFADLTVVY	1674	173		119	5281	133	7012		155
RE DFAFADLTVVYRE	1675	3293		141	4948	60	1728		322
GN TVVYREGNPFGICK	<u>1676</u>	168		121	1833	>13089.91	10064		2407
L GNPFGICKLCLRFL	1677	189		1227	2073	377	13916		45631
S NYSVYGNTLEQTV	1678	14059		1933	91506	822	>14602.8		47481
KK KKPLNEILIRCIICQ	1679 1680	1363		315	1070	347	7972		13328
NEILIRCIICQRPLC	1681	7945		11739	23082	7704	16901		26483
IRCIICQRPLCPQEK	1682	7549		5960	23092	2973	>14602.8		40269
CIVYRDCIAYAACH		1166		928	8560	3973	>14602.8		10186
K NTELYNLLIRCLRC	<u>1683</u>	1108		1366	1293	873	>14602.8		12528
Q IRCLRCQKPLNPAE	1684	7012		6668	9890	8982	>14602.8		>32271.94
K REVYKFLFTDLRIV	1685	8.7	23	112	738	52	54		204
Y RIVYRDNNPYGVCI	<u>1686</u>	524	325	20	432	2307	8307		24147
M NNPYGVCIMCLRFL	1687	1075		1378	2522	454	12020		30895
S	<u>1688</u>								

			HL	A-DR SUP	RTYPE				
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
EERVKKPLSEITIRC	1689	1286		11896	9772	1470	9454		19968
IRCIICQTPLCPEEK	1690	10847		12270	3812	1407	25186		28062
EIPLIDLRLSCVYCK	1691	7610		1876	5012	336	10468		1961
SCVYCKKELTRAE VY	1692	6466		2411	7510	465	8446		2010
VCLLFYSKVRKYR		960	276	286	987	73	258		1798
YY YYDYSVYGATLESI T	1693 1694	1008		186	9855	230	744		1403
IRCYRCQSPLTPEE		10947		13358	83166	10327	13356		>36023.05
K VYDFVFADLRIVYR D	1695 1696	98	2.2	475	5856	717	5962		198
DFVFADLRIVYRDG		6699		867	7197	133	9847		1962
N RIVYRDGNPFAVC	1697	116	144	19	209	1812	6638		4962
KV GNPFAVCKVCLRL LS	1698 1699	134	3805	322	522	56	1034		29300
KKCLNEILIRCIICQ	1700	9357		424	1229	365	16288		3997
NEILIRCIICQRPLC	1701	10992		14069	9339	4621	18947		22062
RTAMFQDPQERPR		9372	154	28192	39014	7977	32947		>25346.4
KL LFVVYRDSIPHAAC H	1702 1703	131	62	3.0	24	690	1998		2855
LTIVYRDDTPHGVC	1704	>15384.62	187	23	203	>8593.4	>72254.34		>25346.4
LCIVYRDCIAYAAC		996	1855	357	1293	628	40121		10660
H YKFLFTDLRIVYRD N	1705 1706	109	8.8	292	256	91	1516		1255
YNFACTELKLVYR		7522	346	1976	4246	3147	2867		2084
DD LKLVYRDDFPYAV	1707 1708	778	237	123	9269	830	28971		18677
CR YDFVFADLRIVYRD G	1709	1160	13	1914	3264	829	21352		5419
LRIVYRDGNPFAVC K	1710	142	181	16	25	557	8985		14207
HEYMLDLQPETTD LY	1711	1377		222	3997	2291	>18559.76		21277
TLRLCVQSTHVDIR T	1712	1517		11996	8650	169	3257		6368
IRTLEDLLMGTLGI	1713	16	5211	95	43	61	895		1718
LEDLLMGTLGIVCP	1714	104		1136	353	1116	261		1994
DLLMGTLGIVCPIC S	1715	966		1324	984	639	963		2614
KATLQDIVLHLEPQ		1204		1987	811	1173	9094		17726
N IDGVNHQHLPARR AE	<u>1716</u> <u>1717</u>	1060		34272	165545	>16971.86	>18559.76		>39914.85
LRAFQQLFLNTLSF	1718	1.5	648	7.4	13	8.3	75		174
FQQLFLNTLSFVCP W		118	1321	134	1585	222	134		2062
W QDYVLDLQPEATD LH	1719 1720	13441		253	45281	5585	>18559.76		>39914.85
DIRILQELLMGSFGI	1721	88	3252	166	290	552	1591		282
IRILQELLMGSFGIV	1722	67	31840	724	710	1208	1998		271
ELLMGSFGIVCPNC		628		1078	8518	1853	4183		949
S KEYVLDLYPEPTDL	1723 1724	5949		131	89674	391	>72254.34		>49867.02
LRTIQQLLMGTVNI V	1724	13	23182	108	208	179	513		181

-			HL	A-DR SUPE	RTYPE				
	SEO ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
Sequence 1QQLLMGTVNIVCP	110.	71	93701	107	483	624	444	0002	156
T QLLMGTVNIVCPTC	1726	1192		2874	10062	4688	2947		2209
A RETLQEIVLHLEPQ	<u>1727</u>	1592		2941	6583	829	25856		19109
N LRTLQQLFLSTLSF	<u>1728</u>	8.3	801	18	18	9.0	60		166
V LQQLFLSTLSFVCP	1729	121	2045	113	754	94	272		152
W KDYILDLQPETTDL	1730	6409		1022	30309	2771	>72254.34		>49867.02
H LRTLQQMLLGTLQ	1731	80	>3750000	437	644	79	6909		5077
VV LQQMLLGTLQVVC	1732	168		1496	631	1068	929		1692
PG QMLLGTLQVVCPG	1733	957		2773	425	3074	3722		2082
CA VPTLQDVVLELTPQ	1734	16056		214	4764	5409	>35360.68		>30612.24
T LQDVVLELTPQTE1	1735	1487		101	1094	417	5673		2180
D QDVVLELTPQTEID	1736	1269		83	1537	53	2716		1684
CKFVVQLDIQSTKE	<u>1737</u>	1251		196	1642	374	4547		19282
D VVQLDIQSTKEDLR	1738	1060		11122	8625	46	3762		13906
DLRVVQQLLMGAL	1739	8.4	25971	325	89	84	508		1845
TV LRVVQQLLMGALT	1740	5.7	21650	115	28	85	82		204
VT VQQLLMGALTVTC	1741	10	34257	239	614	116	71		180
PL QQLLMGALTVTCP	1742	75		1142	1286	201	743		1170
LC QLLMGALTVTCPL	1743	54	>3750000	595	870	1019	389		303
CA REYILDLHPEPTDL F	1744	154		132	9957	354	7257		29316
TCCYTCGTTVRLC1	1745	1230	19884	719	2269	132	63		1374
N VRTLQQLLMGTCTI V	<u>1746</u> 1747	36	32360	322	39	114	1820		496
LQQLLMGTCTIVCP		197		1147	483	522	2098		1638
MLDLQPETTDLYC YE	1748 1749	10076	720	1913	12241	4249	>72254.34		>32230.34
VLDLYPEPTDLYCY E	1750	11201	121	203	2193	212	>72254.34		>32230.34
LREYILDLHPEPTD	1751	134	891	23	9235	968	21989		16462
HIEFTPTRTDTYAC	1752	50000	30000	667	10000		>12500		
RV LWWVNNESLPVSP RL	1752		315						
YEEYVRFDSDVGE	1754	50000		400	500000		250000		
EEYVRFDSDVGE	1755	50000		216	500000		250000		***
APPRLICDSRVLER Y	1756	1374	6.3	9735	5794	7141	8937	11214	9348
ICDSRVLERYLLEA K	1757	2758	236	1984	10984	11016	57605	808	>78947.37
VLERYLLEAKEAE NI	<u>1758</u>	933	59010	2598	12139	5019	13067	3150	6382
EHCSLNENITVPDT K	1759	9837	27481	2294	28297	1205	32375	6191	>78947.37
NENITVPDTKVNFY A VPDTKVNFYAWKR	1760	>24154.59	4.8 259	>21390.3 7 1742	7612 4131	>18572.83 1328	42846 38622	1850 422	>78947.37 >78947.37
ME	<u>1761</u>	2704	239	1742	4131	1320	30022	422	~10941.31

HLA-DR SUPERTYPE										
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901	
VNFYAWKRMEVG		193	2871	10	291	15	40163	35	1238	
QQA WKRMEVGQQAVE VWQ	1762 1763	62	514	24	2591	94	46062	139	14696	
VGQQAVEVWQGL		161	>174081.24	10294	6283	923	4230	>40511.09	>78947.37	
ALL VEVWQGLALLSEA	1764	86	13293	1310	1357	79	6863	13411	8151	
VL GLALLSEAVLRGQ	1765	83	816	11	21	1435	4606	2000	15148	
AL SEAVLRGQALLVN	<u>1766</u>	11	70855	2064	4207	17446	1087	>63636.36	>78947.37	
SS RGQALLVNSSQPW	<u>1767</u>	1118	93874	1697	1168	3434	319	29454	8450	
EP LVNSSQPWEPLQL	1768	2178	26138	>21505.3	13031	19689	8344	16920	>78947.37	
HV QPWEPLQLHVDKA	1769	11567	4862	8 1296	6135	1111	24157	>63636.36	34819	
VS LQLHVDKAVSGLR	<u>1770</u>	192	22	9.7	44	13571	3213	801	>78947.37	
SL DKAVSGLRSLTTLL	1771	13	4331	1014	25	247	615	16375	>78947.37	
R GLRSLTTLLRALGA	<u>1772</u>	8.5	2345	24	9.2	30	509	14	1136	
Q TTLLRALGAQKEA1	<u>1773</u>	19	107164	339	199	103	4281	652	4607	
S ALGAQKEAISPPDA	<u>1774</u>	194	>204081.63		93062	13015	>71225.07	>60214.56	15337	
A KEAISPPDAASAAP	<u>1775</u>	15531	48560	8 6590	4389	28755	6661	6391	5735	
L PPDAASAAPLRTIT	<u>1776</u> .	309	14900	566	68	1555	24937	>63636.36	8674	
A SAAPLRTITADTFR	<u>1777</u>	1166	1262	1185	261	1456	3646	28110	2505	
K RTITADTFRKLFRV	<u>1778</u>	148	139	1042	928	1957	3448	792	4692	
Y DTFRKLFRVYSNFL	<u>1779</u>	12	6946	70	104	93	10	39	307	
R LFRVYSNFLRGKLK	1780	43	6156	643	1816	1275	5.5	28	3508	
L SNFLRGKLKLYTGE	<u>1781</u>	143	9583	2883	2375	7182	3783	1433	8099	
A KLKLYTGEACRTG	<u>1782</u>	122	18435	5964	3505	36294	8082	7683	2860	
DR APPRLITDSRVLER	1783	10144	15	6680	3168	7765	629	26382	8391	
Y ITDSRVLERYLLEA	<u>1784</u>	1571	6501	1303	1990	13339	7498	967	>78947.37	
K EHTSLNENITVPDT	<u>1785</u>	43921	33635	12379	2769	1245	37154	>16333.33	>78947.37	
K KLKLYTGEATRTG	1786	178	118459	15	3230	1426	8234	2008	>78947.37	
DR PQPFRPQQPYPQ	<u>1787</u>									
PFRPQQPYPQ	<u>1788</u>									
	1789									
PQPFRPQQPYP	<u>1790</u>									
PQPFRPQQP	<u>1791</u>									
KOPFRPOOPYPO										

KQPFRPQQPYPQ

PKPFRPQQPYPQ

PQPFKPQQPYPQ

PQPFRKQQPYPQ

PQPFRPQKPYPQ

PQPFRPQQPKPQ

PQPFRPQQPYKQ

PQPFRPQQPYPK

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HLA-DR SUPERTYPE										
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901	
QFLGQQQPFPPQ	1800									
FLGQQQPFPPQ										
LGQQQPFPPQ	1801 1802									
QFLGQQQPFPP	1803									
QFLGQQQPF										
IRNLALQTLPAMCN	<u>1804</u>									
VY NLALQTLPAMCNV Y	1805 1806									
LALQTLPAMCNVY	1807									
IRNLALQTLPAM	1808									
IRNLALQTLP	1809									
EGDAFELTVSCQG	1009			572	3578					
GLPK ESTGMTPEKVPVSE VMGT	1810 1811	>50000	>47368.42	510	>71428.57		>31250			
FPTIPLSRLFDNASL	1812	8071	114611	228	22	7210	3175	4969	9876	
RLFDNASLRAHRL		89	97	77	2043	10328	1921	14985	23832	
HQ LRAHRLHQLAFDT YO	1813 1814	162	15603	5076	2197	10139	123	5621	15115	
QLAFDTYQEFEEA		>20491.8	7981	>10738.2	33446	5399	2580	>33333.33	>59523.81	
YI QEFEEAYIPKEQKY S	1815 1816	>20491.8	>171755.73	6 >21276.6	>88339.22	395	31344	>33333.33	>59523.81	
IPKEQKYSFLQNPQ		128	49978	217	3633	9.0	8305	13553	79800	
T SFLONPOTSLCFSES	<u>1817</u>	595	8617	6376	16880	>25832.77	48620	>33333.33	93856	
TSLCFSESIPTPSNR	1818	604	182762	48	229	852	1064	>33333.33	4395	
REETQQKSNLELLR	<u>1819</u>	8921	91054	9341	1324	1433	51179	22467	9680	
I SNLELLRISLLLIQS	<u>1820</u>	72	43487	621	189	379	642	>33333.33	3422	
ISLLLIQSWLEPVQF	<u>1821</u>	184	27922	885	177	0.86	83	>33333.33	6247	
SWLEPVQFLRSVFA	1822	11	167103	1128	152	883	589	3416	3998	
N	1823									
FLRSVFANSLVYGA S	1824	4.3	15221	6.7	43	59	16	13436	15127	
NSLVYGASDSNVY		7313	81158	190	1585	1055	201	>33333.33	3896	
DL SDSNVYDLLKDLE	1825	24369	54982	11032	>25680.53	95	182355	>33333.33	>59523.81	
EG GIQTLMGRLEDGSP R	1826 1827	98	>55900.62	11914	2458	3745	18952	>33333.33	37821	
RLEDGSPRTGQIFK		15693	76675	7906	1729	22125	35120	>33333.33	>59523.81	
Q RTGQIFKQTYSKFD T	1828 1829	1555	20341	1680	1831	40	46	16432	8515	
QTYSKFDTNSHND		17352	>55900.62	97	11218	78	54569	7726	31341	
DA TNSHNDDALLKNY GL	1830 1831	16457	26397	20308	>25680.53	16329	245523	>33333.33	>59523.81	
ALLKNYGLLYCFR		137	9819	446	1286	551	11915	>33333.33	676	
KD DMDKVETFLRIVQ CR	1832 1833	1277	4813	867	1135	622	10484	1673	16127	
FLRIVQCRSVEGSC		106	33536	185	164	191	7199	7262	5311	
GF FPTIPLSRLFDNAM L	1834 1835	6923	46707	9458	175	923	5529	1051	14964	
RLFDNAMLRAHRL		2.3	27	6289	1520	4247	3297	212	>59523.81	
HQ QLAFDTYQEFEQNP Q	1836 1837	>17985.61	7851	28586	47399	4843	21064	>33333.33	>59523.81	

HLA-DR SUPERTYPE									
Sequence	SEO ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
SFLQNPQTSLCCFR		106	1829	671	1816	1230	7026	7069	3082
K SNLELLRICLLLIQS	1838	731	61913	1526	2303	1112	1222	19782	3970
ICLLLIQSWLEPVQF	1839	8511	50874	11303	5708	71	643	>33333.33	>59523.81
NSLVYGASDSNIYD	1840	13068	>51428.57	240	3683	1229	297	>33333.33	>59523.81
L SDSNIYDLLKDLEE	1841	>17985.61	124500	17458	25922	137	>85034.01	>33333.33	50134
G DKVETFLRIVQCCG F	1842 1843	953	18325	1158	259	397	697	581	4080
SFLQNPQTSLTFSES	1844	1191	2395	7780	15527	9558	6197	>33333.33	17714
TSLTFSESIPTPSNR	1845	182	17425	18	98	686	682	17602	2461
ALLKNYGLLYTFR		19	5982	160	266	303	5923	3616	2628
KD LLYTFRKDMDKVE TF	1846 1847	>17985.61	23871	10623	17771	1133	53362	10448	>59523.81
DMDKVETFLRIVQ TR	1848	1111	11194	2030	133	454	436	183	51511
FLRIVQTRSVEGST	_	6.4	3944	11	16	99	9.8	445	778
GF HLDMLRHLYQGCQ VV	1849 1850	304	37552	9417	2741	3593	27027	5384	12508
RLRIVRGTQLFEDN		4.8	11287	8389	2929	1024	12	6325	1834
YAL GVGSPYVSRLLGIC	1851	19	167949	1570	49	4156	190	1317	2614
TLERPKTLSPGKNG	1852 1853	10103	134367	>22471.9	103285	>28592.93	25988	>75384.62	>300000
KIFGSLAFLPESFDG		597	74162	1195	1897	37	377	>75384.62	15796
DPA ELVSEFSRMARDPQ	1854 1855	201	1026	120	4882	15120	21259	4082	91575
GEALSTLVLNRLK		719	11783	3045	305	14802	3191	192	20167
VG AYVLLSEKKISSIQS	1856 1857	78	136	943	359	9471	3848	27	3338
VASLLTTAEVVVTE	1858	604	136308	7431	810	6517	369	>118357.49	1955
KCEFQDAYVILLSE	1859	14	5791	73	943	351	336	489	185
KK ALSTLVLNRLKVG		49	153	517	31	2167	647	4.0	2166
LQ MSYNLLGFLQRSS	1860	115	156715	366	1584	788	1060	3421	3646
NC LGFLQRSSNCQCQ	1861	437	112406	120	401	827	767	218	3729
KL RSSNCQCQKLLWQ LN	1862	9665	>191897.65	1046	2987	12652	9689	4530	74405
QCQKLLWQLNGRL	1863	181	133472	360	460	1004	3702	2519	4669
EY LWQLNGRLEYCLK	1864	1108	2356	816	8882	1024	10586	>16333.33	5206
DR GRLEYCLKDRRNF DI	1865 1866	9854	853	918	4155	3238	12108	1318	25159
RNFDIPEEIKQLQQF	1867	6969	26262	18107	5375	>114457.83	47893	>144117.65	>77319.59
PEEIKQLQQFQKED		1026	40154	1618	618	7875	49505	11908	>77319.59
A QLQQFQKEDAAVT IY	1868 1869	85	17383	231	27473	1121	500	4862	55351
QKEDAAVTIYEML QN	1859	8376	>156521.74	9437	75877	785	45455	>144117.65	5989
AVTIYEMLQNIFAIF	1871	17	23730	101	808	163	267	6873	4540
EMLQNIFAIFRQDS S	1872	395	9544	685	689	456	3313	10429	9738
IFAIFRQDSSSTGW N	1873	132	402	9.6	71	118	1186	4725	970
RQDSSSTGWNETIV E	1874	>102040.8	38681	4637	184507	40847	36320	15135	9075
-	1017	-							

HLA-DR SUPERTYPE										
Sequence	SEQ ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901	
STGWNETIVENLLA		21407	>156521.74	1755	10422	7060	3960	>144117.65	>77319.59	
N ETIVENLLANVYHQ R	1875 1876	659	40053	789	802	326	21681	>144117.65	8151	
NLLANVYHQRNHL		152	40328	1039	1440	1492	8000	453	4160	
KT VYHQRNHLKTVLE	<u>1877</u>	617	3135	7757	76003	153	6180	2101	>77319.59	
EK LEKEDFTRGKRMS SL	<u>1878</u> 1879	21965	50733	>20887.7	93968	5694	946	804	>77319.59	
FTRGKRMSSLHLK RY	1880	13	3302	1013	970	484	136	553	10925	
RMSSLHLKRYYGRI L	1881	275	2181	993	4793	34	283	277	14964	
HLKRYYGRILHYL KA	1882	26	3709	135	666	86	214	237	2896	
YGRILHYLKAKEDS H	1883	30	42429	2343	917	23	900	704	7577	
HYLKAKEDSHCAW TI	1884	1128	34758	2064	12153	3701	581	34851	>77319.59	
KEDSHCAWTIVRV	1885	4835	>46656.3	353	1090	74	30	40000	2937	
CAWTIVRVEILRNF Y	1886	66	3561	158	640	135	746	43672	757	
VRVEILRNFYVINR	1887	1.8	429	140	47	18	14	3585	485	
RNFYVINRLTGYLR N	1888	1.7	2199	219	4618	182	527	167	7600	
MSYNLLGFLQRSS NT	1889	25	107838	1152	813	433	8867	900	8972	
LGFLQRSSNTQTQK L	1890	142	26455	18	211	1068	420	939	1345	
RSSNTQTQKLLWQ LN	1891	10515	44338	2139	15497	12590	27678	1283	>77319.59	
QTQKLLWQLNGRL EY	1892	32	3555	55	35283	86	3099	2042	2083	
LWQLNGRLEYTLK DR	1893	698	511	757	16171	94	20198	43286	16619	
GRLEYTLKDRRNF DI	1894	7252	30	3228	97035	1379	4961	4917	>77319.59	
HYLKAKEDSHTAW TI	1895	232	70237	553	10677	15067	801	8526	10140	
KEDSHTAWTIVRV EI	1896	1909	44754	746	2178	302	35	>79032.26	6079	
TAWTIVRVEILRNF Y	1897	7.8	2997	44	84	115	29	57243	404	
LGFLQRSSNCQSQK L	1898	192	4888	8.1	93	228	305	405	13167	
RSSNCQSQKLLWQ LN	1899	2050	57946	595	16721	4010	8922	6943	4062	
QSQKLLWQLNGRL EY	1900	127	33374	84	741	55	1166	991	5920	
GIVEQCCTSICSLY Q	1901	11123	777105	10911	2995	17793	>79872.2	>10047.16	13855	
TSICSLYQLENYCN	1902	11391	>154109.59	20462	3791	12457	>85616.44	>54444.44	>63025.21	
GILEQCCTSICSLYQ	1903	11025	>187500	14862	5106	15983	54113	>54444.44	16714	
GIVEQTTTSITSLYQ EQTTTSITSLYQLE	<u>1904</u>	6354 18953	107486 >143769.97	121 170	115 258	818 272	788 2230	>54444.44 >54444.44	13304 17381	
N TSICSLYQLENYCG	1905	1125	202253	8841	1986	1089	247525	>54444.44	>83333.33	
TSITSLYQLENYTN	1906	1253	81293	1468	138	851	6055	26791	9947	
TSITSLYQLENYTG	1907	1132	96727	1628	129	115	8371	14562	46268	
GIVEQCCCGSHLVE	<u>1908</u>	10043	>74750.83	19904	2892	6626	41276	>54444.44	>63025.21	
A SLYQLENYCCGER	1909	3568	54469	7313	1527	2356	12308	>54444.44	>83333.33	
GF CCTSICSLYQLENY	<u>1910</u>	11655	71239	8383	1604	629	35604	>54444.44	29845	
CC	<u>1911</u>									

-			HL.	A-DR SUPE	RTYPE				
Sequence	SEQ ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
GSHLVEALYLVCC N	1010	194	>59681.7	2280	11512	2509	302	>54444.44	37166
CCGSHLVEALYLV CC	<u>1912</u> <u>1913</u>	880	>55693.07	10081	20487	5230	1822	>54444.44	>63025.21
FVNQHLCGSHLVE		583	>187500	19209	39746	>20663.4	6791	>54444.44	>63025.21
AL QHLCGSHLVEALY LV	<u>1914</u> 1915	170	48557	12954	4303	9825	86	>54444.44	7422
GSHLVEALYLVCG		525	>187500	8292	1603	4609	560	>54444.44	5386
ER VEALYLVCGERGF FY	<u>1916</u> 1917	76	17558	209	124	1044	3869	24623	2233
YLVCGERGFFYTPK T	1918	11063	37210	1439	22980	730	64644	>54444.44	1520
FVNQHLCGSDLVE		117	>74750.83	19154	36693	14913	38662	>54444.44	>63025.21
AL FVNQHLTGSHLVE	<u>1919</u> 1920	9.2	67240	858	14916	1065	15	>54444.44	41482
AL QHLTGSHLVEALY		9.3	50338	>16096.5	3952	7423	38	>54444.44	42312
LV GSHLVEALYLVTG ER	1921	645	>176470.59	15781	1693	14443	553	>54444.44	>63025.21
VEALYLVCGERGS FY	1922 1923	88	9972	833	194	6108	6485	>54444.44	6311
VEALYLVCGERGF LY	1924	14	11587	167	31	1027	5351	10565	3063
VEALYLVTGERGFF Y	1925	9.9	2011	60	23	2342	195	1224	683
YLVCGERGFLYTP KT	1926	155	2033	>20460.3	>38550.5	>30134.81	12842	>54444.44	124
YLVCGERGFFYTD KT	1927	17260	11790	>20460.3	>38550.5	>30134.81	92272	>54444.44	317
YLVCGERGFFYTKP T	1928	3207	42139	>20460.3	>38550.5	>30134.81	969	>54444.44	1673
YLVTGERGFFYTPK T	1929	779	517	>20460.3	>38550.5	30457	7737	29236	6295
YLVTGERGFFYTD KT	1930	3259	7326	>20460.3	>38550.5	>30134.81	5328	>25789.47	2909
YLVTGERGFFYTKP T	1931	1152	4801	>20460.3	>38550.5	>30134.81	78	4304	195313
VCGERGFFYTPKTR R	1932	9622	1989	>20460.3	>38550.5	>15103.34	5494	419	14379
VTGERGFFYTPKTR R	1933	18906	3018	7226	147000	13417	27824	9407	>300000
MWDLVLSIALSVG CT	1934	205		1846			3032	23046	1727
DLVLSIALSVGCTG	1935	1197		13038			4029	>245000	2200
HPQWVLTAAHCLK KN	1936	22	1103	875			563	1693	822
QWVLTAAHCLKK NSQ	1937	895		>40000			3402	98000	4813
GORVPVSHSFPHPL	1938	1563		>40000			629	>245000	102
RVPVSHSFPHPLYN M	1939	67		>16000			101	100021	97
PHPLYNMSLLKHQ SL	1940	19079		819			20691	3315	1592
HPLYNMSLLKHQS LR	1941	232	13007	499			1282	382	199
NMSLLKHQSLRPD ED	1942	3131		>40000			20620	26496	96825
SHDLMLLRLSEPAK I	1943	56	2396	2244			106	1327	112
HDLMLLRLSEPAKI T	1944	16	1406	3063			109	544	43
PEEFLRPRSLQCVS L	1945	2001		>26666.6 7			5156	2207	5839
PRSLQCVSLHLLSN D	1946	1111		16000			2217	6107	28307

	HLA-DR SUPERTYPE										
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901		
NGVLQGITSWGPEP		1093		8433			2285	52234	50111		
C KPAVYTKVVHYRK WI	<u>1947</u> 1948	5000		1433			2401	53	3677		
LHLLSNDMCARAY		2104	938	4277			27685	50230	59904		
SE VGNWQYFFPVIFSK	1949	37		4.1			100				
A ESEFQAALSRKVA KI.	<u>1950</u> 1951			579	29617						
IGHLYIFATCLGLS				>816.33	12199						
YDGL VGNWQYFFPVIFSK ASDSLQLVFGIELM	<u>1952</u>			654	3846						
EVD PAYEKLSAEQSPPP V	1953			479	>250000						
RNGYRALMDKSLH	1954			512	5779						
VGTQCALTRR FFKNIVTFFKNIVT	<u>1955</u>	50000		>666.67	500000		>12500				
YKSAHKGFKGVDA	<u>1956</u>	70	>900000	889	25000		108				
QGTLSKI VDAQGTLSKIFKLG	1957	25	1383	1600	314		1171				
GRDSRS AC-	<u>1958</u>	50000	>900000	889	25000		2362				
ASQKRPSQRHGSK YLATAST ENPVVHFFKNIVTP	<u>1959</u>										
R ENPVVAFFKNIVTP	1960										
R ENPVVHAFKNIVTP	<u>1961</u>										
R ENPVVHFFANIVTP	1962										
R ENPVVHFFKNIVTP	1963										
A NPVVHFFKNIVT	1964										
HFFKNIVTPRTPPY	<u>1965</u> 1966										
NPVVHFFKNIVTPR	1967										
LPVPGVLLKEFTVS	1968	57	15058	14	12	12	57				
GNILTI WITQCFLPVFLAQP PSGQRR	1969	679	25534	88	2804	216	74162				
DHRQLQLSISSCLQ		1356	42666	1322	210	725	736				
QLSLLM YLAMPFATPMEAE	<u>1970</u>	46	46591	266	814	405	526				
LARRSLA AAPLLLARAASLSL G	1971	6.8	35410	139			160	30	64		
APLLLARAASLSLG	1972	8.4	56250	202			59	76	124		
F PLLLARAASLSLGF	1973	10	>81818.18	521			162	37	58		
L SLSLGFLFLLFFWL	1974	11417		4711			22727	>122500	24620		
D LLFFWLDRSVLAK	1975	2.9	6.3	2.6			135	163	518		
EL DRSVLAKELKFVTL	1976	705		569			2016	15815	4719		
V AKELKFVTLVFRH	1977	787	30000	783			606	1953	2355		
GD RSPIDTFPTDPIKES	1978 1979	>50000		13095			>62500	>245000	6124		
FGQLTQLGMEQHY		2259		3210			>62500	109567	>187500		
EL DRTLMSAMTNLAA	1980	97	64286	13			383	2362	222		
LF MSAMTNLAALFPP	1981 1982	1757		700			36084	73870	>187500		

			HL	A-DR SUPE	RTYPE				
Sequence	SEQ ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
EG				- 40000			- 125000	39231	22022
MTNLAALFPPEGVS 1	1983	24		>40000			>125000		22822
PEGVSIWNPILLWQ P	1984	111		1778			15030	28577	103096
GVSIWNPILLWQPI		44	56250	10328			4992	11008	3985
P WNPILLWQPIPVHT	<u>1985</u>	208	>81818.18	695			521	115494	607
V NPILLWQPIPVHTV	<u>1986</u>	31	>81818.18	206			41	12999	575
P PILLWQPIPVHTVPL	1987	44	>81818.18	258			46	21244	168
ILLWOPIPVHTVPLS	1988	45	>81818.18	170			19	13091	131
WQPIPVHTVPLSED	<u>1989</u>	6386		>26666.6			159	>81666.67	17518
Q LSGLHGQDLFGIWS	1990	148		7 >26666.6			>35714.29	>81666.67	>125000
K YDPLYCESVHNFTL	<u>1991</u>	1597	16625	7 8889			838	30867	643
P	1992		10023						
LPSWATEDTMTKL RE	1993	20274		973			>35714.29	>81666.67	>125000
LRELSELSLLSLYGI	1994	655		371			4010	9368	1614
LSELSLLSLYGIHK O	1995	482	>81818.18	1549			20906	1186	1450
LSLLSLYGIHKQKE K	1996	656	>81818.18	4444			>35714.29	1637	4959
KSRLQGGVLVNEIL		362		>26666.6			2838	>81666.67	5516
N GGVLVNEILNHMK	<u>1997</u>	2165	700	359			29463	3239	54411
RA IPSYKKLIMYSAHD	<u>1998</u>	9.9	9728	510			1946	60	351
T YKKLIMYSAHDTT	1999	17	22678	207			292	309	107
VS	2000	4496	22010	24			731	24812	813
LIMYSAHDTTVSGL Q	2001								
DTTVSGLQMALDV YN	2002	171		4424			14706	>245000	2876
ALDVYNGLLPPYA SC	2003	18		485			>83333.33	588	86603
LDVYNGLLPPYAS		15		348			>83333.33	404	31277
CH YNGLLPPYASCHLT	2004	42		6189			>83333.33	14027	8022
E FAELVGPVIPQDWS	2005	12		4690			24056	>245000	39472
T TVPLSEDQLLYLPF	2006	4012	332	10755			11313	42162	37369
R	2007	2249	592	8051			13062	18841	26949
LTELYFEKGEYFVE M	2008	2249		8031			13002	10041	20949
GPVIPQDWSTECM TT	2009		52098						
QAHSLERVCHCLG KWLGHPDK	2010	50000		667	500000		>250000		
WTTCQSIAFPSKTS			17308	22					
ASIGSL QKGRGYRGQHQA	2011		>47368.42	88					
HSLERVCH AATYNFAVLKLMG	2012		>52941.18	533					
RGTKF VATGLCFFGVALFC	<u>2013</u>		>112500	351					
GCGHEA	2014								
FLYGALLLAEGFYT TGAVRQ	2015								
SAVPVYIYFNTWTT									
CQSIAF TLSVTWIGAAPLIL	2016	3.1	>81818.18	7273			16	840	5.4

	HLA-DR SUPERTYPE										
Sequence	SEO ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901		
SVTWIGAAPLILSRI	2018	4.1	>81818.18	3152			83	139	30		
VTWIGAAPLILSRIV	2019	8.1	>81818.18	8000			195	731	82		
SQPWQVLVASRGR		66	>81818.18	7628			385	386	621		
AV GRAVCGGVLVHPQ	2020	386		>26666.6			3582	>245000	8069		
WV GVLVHPQWVLTAA	2021	87	21320	7 67			153	1931	365		
HC HPQWVLTAAHCIR	2022	13	3632	1621			283	1305	107		
NK QWVLTAAHCIRNK	2023	50		19403			214	2598	967		
SV AHCIRNKSVILLGR	2024	578	29704	69			2573	104	715		
H SVILLGRHSLFHPE	2025	717	1400	12649			26088	500	5216		
D VILLGRHSLFHPED	2026	273	8744	8208			30625	737	18520		
T GQVFQVSHSFPHPL	2027	288	45000	8.2			27	548	33		
Y VFQVSHSFPHPLYD	2028	16	>75000	25			51	8751	17		
M PHPLYDMSLLKNR	2029	1315		20787			10699	29813	12836		
FL SHDLMLLRLSEPAE	2030	532	6215	4051			58	3538	64		
L HDLMLLRLSEPAEL T	2031	62	2867	6193			152	3914	22		
TDAVKVMDLPTQE PA	2032 2033	>50000		>80000			>41666.67	20875	>107142.8		
LHVISNDVCAQVH PO	2033	789	8318	790			17451	>122500	32671		
CAQVHPQKVTKFM LC	2035	10206		2566			32275	8731	34893		
GGPLVCNGVLQGIT	2036	3353		68			>35714.29	9334	16308		
GPLVCNGVLQGITS W	2037	1724		30			4893	4187	32640		
NGVLQGITSWGSEP C	2038	945	24942	560			485	5874	819		
RPSLYTKVVHYRK WI	2039	6041	53785	339			652	39	5484		
HSLFHPEDTGQVFQ V	2040		65260								
PRWLCAGALVLAG GF	2041	46		>20000			766	26531	1439		
LGFLFGWFIKSSNE A	2042	10	>75000	1338			2261	1421	1701		
LDELKAENIKKFLY N	2043	1136	1370	4842			7470	1248	12778		
IKKFLYNFTQIPHL A	2044	449	8080	43			29	512	160		
KFLYNFTQIPHLAG T	2045	340	13805	217			30	415	54		
WKEFGLDSVELAH YD	2046	1139	85	96			3511	19971	7052		
LAHYDVLLSYPNK TH	2047	79	37533	1117			3617	415	1009		
GNEIFNTSLFEPPPP GKVFRGNKVKNAO	2048	20412 612		>20000 1087			>35714.29 2350	>163333.33 4121	10415 31277		
LA GNKVKNAQLAGA	2049	677		13333			>83333.33	28904	7882		
KGV EYAYRRGIAEAVG	2050	5.1		213			70	596	67		
LP	2051			9923			2015	>490000	23102		
AEAVGLPSIPVHPIG AVGLPSIPVHPIGY	2052	5.4 3.6		4193			1080	4432	15377		
Y	2053										

			HI	A-DR SUPE	RTYPE				
Sequence	SEQ ID NO.	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
IGYYDAQKLLEKM		1923		12649			>83333.33	8236	47246
GG TGNFSTQKVKMHI HS	2054 2055	11180		833			9407	10282	1450
TRIYNVIGTLRGAV E	2056	14	33333	6.3			4806	70	2900
ERGVAYINADSSIE G	2057	2440		6761			34021	>163333.33	25516
GVAYINADSSIEGN Y	2058	1054		146			6244	23360	3048
DSSIEGNYTLRVDC T	2059	16667		3360			14458	>163333.33	>187500
NYTLRVDCTPLMY SL	2060	6804	45	9.9			24597	6323	48412
CTPLMYSLVHNLT KE	2061	93	19437	245			140	223	249
DFEVFFQRLGIASG R	2062	143		221			21926	122	2005
EVFFQRLGIASGRA R	2063	28	>75000	22			5311	6.3	2976
TNKFSGYPLYHSV YE	2064	3402		5521			30853	614	741
YDPMFKYHLTVAQ VR	2065	9.0	>75000	19			158	172	179
DPMFKYHLTVAQV RG	2066	5.7	>75000	9.1			168	43	258
MFKYHLTVAQVRG GM	2067	16	29032	18			72	70	266
KYHLTVAQVRGG MVF	2068	137	33658	806			228	1519	5860
VAQVRGGMVFELA NS	2069	228		662			4449	>98000	499
RGGMVFELANSIVL P	2070	10	37118	229			41	8682	33
GMVFELANSIVLPF D	2071	15	4604	230			30	4995	81
VFELANSIVLPFDC R	2072	19	667	999			39	36123	50
ADKIYSISMKHPQE M	2073	22361		5310			4098	1136	3512
IYSISMKHPQEMKT Y	2074	8452		16000			11573	1357	12293
PQEMKTYSVSFDSL F	2075	15143		3024			1192	>98000	1981
TYSVSFDSLFSAVK N	2076	219	101	73			346	2256	526
VLRMMNDQLMFL ERA	2077	118	183	29			17334	1700	10684
LRMMNDQLMFLER AF	2078	2704		392			17507	2492	4601
RHVIYAPSSHNKYA G	2079	2174		481			31250	11667	481
RQIYVAAFTVQAA AE	2080	3.7	28347	1.2			292	36	91
QIYVAAFTVQAAA ET	2081	1.6	26609	1.6			324	102	65
VAAFTVQAAAETL SE	2082	14	>75000	58			793	1420	127
YISIINEDGNEIFNT	2083	498	397	624			23719	>122500	83056
ISINEDGNEIFNTS	2084	507	559	>12965.9 6 226			>23105.36 8550	>122500	>52337.75 >52337.75
EDFFKLERDMKINC S FFKLERDMKINCSG	2085	2710 4419	468 121	483			>23105.36	8109	>52337.75
K GVILYSDPADYFAP	2086	1566	17	7508			>23105.36 7848	106291	>52337.75 2473
G GAAVVHEIVRSFGT	2087	1300	17	/308			/040	100291	24/3
L NSRLLQERGVAYIN	2088 2089	614	318	5089			7997	3224	2616

A VAYINADSSIEGNY		HLA-DR SUPERTYPE										
VATINADSSIEGNY	Sequence											
TODQIMFLERAFIDPL G	A											
DOLIMERRAFIDPL Sept		2090	4716	531	411			9745	105832	5467		
SANPLICYYSGFHP 2092	DQLMFLERAFIDPL			>19667.83								
ACP MPDAENWNSQFEIL EDAA 2093 50000 22500000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 2250000 22500000 22500000 2250000000000	KSNFLNCYVSGFHP		2500	>900000	296	3125		8333				
EDAM SEVILISARDVLAVV S S	AC-	2092					500000		>25000			
EVILISARDVLAVV S 1	NPDAENWNSQFEIL	2093										
YKTIAYDEEARR 2095 50000 143 4000 500000 250000 250000 250000 27 PYILLVSKKING 0 2096 1.1 106 64 13 136 38 12 134 140 804 804 804 805 805 806 805 807 806 808 807 808 808 808 808 808 808 808 808	EYLILSARDVLAVV				508							
GEALSTLVYNKIRG			50000	143	4000	500000		250000				
PYILLYSKYSTYK			292	29687	1535	246	30057	2325	383	40840		
EAVLEDPYILLVSS 34	PYILLVSSKVSTVK		1.1	106	64	13	136	38	12	134		
AGLIFTITEAVVAD Company	EAVLEDPYILLVSS		34	479	233	172	681	933	1666	15032		
X		2098	6.8	27189	13	106	67	230	3893	409		
X	K	2099	75	274	648	40	3626	396	20	18035		
SELISTNLPYGRTN 2102 4226 690 50000	K	2100										
L C 102		2101					3363		13336			
March 1160 283 4868 10869 1161 1160 283 4868 10869 1161 1161 1210 1161 1210	L	2102							10641			
LIFLYKMNALRRLP V V V 2105 2105 2106 2107 2106 2107 2107 2108 2107 2108 2109 2109 2109 2109 2109 2109 2109 2109	M	2103										
V V V V V V V V V V	L	2104										
MNALRIPYICSFL 2.1 13 488 265 SAFLESQSMNKIGD 549 113 523 21493 LKELIKVGLPSFEN 1 2109 FENLVAENVEPK 2109 PATYGINPVLTSLENK V 2111 LIKIIWKNYMKIMN HL 121 132 MTLYQIQVMKRNQ KQ QQVQMMIMIKFM OMINKFMOVINII 2115 2115 2115 2115 2115 2115 2115 21		2105	0.80		5.6			56	19			
SAFLESQSMNKIGD D C DE CONTROLLE CONT	MNALRRLPVICSFL		2.1		13			488	265			
LKELIKVGLPSFEN L LKELIKVGLPSFEN L 2108 99 163 542 1493 L FENLVABNYKPK 2109 56 2372 120215 >25025.54 V VD 2109 1.03 15 139 181 V PATYGINPVLTSLENK V V 2111			549		113			523	21493			
FENLYAENVKPPK VD	LKELIKVGLPSFEN		99		163			542	1493			
PATYGIIVPVLTSLFY ZIII0 1.03 15 139 181 YGIIVPVLTSLFNK V ZIIIV 1.03 15 130 15 139 181 YGIIVPVLTSLFNK V ZIIIV 1.03 15 130 15 130 15 130 15 130 15 130 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 130 15 15 15 15 15 15 15 15 15 15 15 15 15	FENLVAENVKPPK		56		2372			120215	>25025.54			
YGIIVPVLTSLEPNK V 2111			1.03		15			139	181			
LLKIWNYMKINN	YGIIVPVLTSLFNK		6.0		2.0			60	793			
MTLYQIQVMKRNQ	V LLKIWKNYMKIMN	2111	121		132			395	132			
NO 10 10 10 10 10 10 10 1	HL	2112	1219					31053	166			
GV 2114 312 68040 66150 MMIKEMGVIYIMII 215 2905 312 68040 66150 GVIYIMIISKKMMR 10 22 476 137 LYYLFNQHIKKELY 216 12 1224 10244 1771 HFNMLKNKMQSSF 12 18 3225 185 FM 2119 2834 1492 >88339.22 1204 K 2119 73 24 11942 13255 NK 212 55 1839 3578 180 ENEYATGAVRPPQ 4438 281 4970 17329 AA NYELSKKAVITFI 713 536 5498 141 Y 2123 993 303 534 2240 CKUKDINERDEN 438 16 4438 1785	KQ	2113							182			
Comparison	GV	2114										
Comparison Com		2115										
HENMILKNKMQSSF FM 2118 2118 2118 2118 2118 2118 2118 211	K	2116										
FM 2118 2834 1492 >88339.22 1204	H	2117										
LDIVQKLYIKQEEQ		2118	12		18			3225	185			
QKKYIYNLIMNTQ 73 24 11942 13255 NK 2120 55 1839 3578 180 YEALIKLLPFSKRIR 2121 55 1839 3578 180 EMEYATGAVRPPQ 4438 281 4970 17329 AAA 2122 713 536 5498 141 VELIKIPYTKNITPI 2123 993 303 534 2240 VELIKIPSKEPEN 438 16 4438 17850	LDIYQKLYIKQEEQ		2834		1492			>88339.22	1204			
YEALIKLIPFSKRIR 2121 55 1839 3578 180 ENEYATGAVRPFQ 4438 281 4970 17329 AA SLOW 1713 536 5498 141 YELIKIPTYKNITI 2123 QKILIKIPTYKNITI 2124 993 303 534 2240 CKILIKIPTYKNITI 2124 678 16 46383 17859	QKKYIYNLIMNTQ		73		24			11942	13255			
ENEYATGAVRPFQ 4438 281 4970 17329 AA 2122 713 536 5498 141 V 2123 QKILKIPYTKNIT 2124 993 303 534 2240 CKILKIPYTKNIT 2124 678 16 46333 17850			55		1839			3578	180			
NYELSKKAVIFTPI 713 536 5498 141 7 2123 9 3 303 534 2240 8 141 9 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			4438		281			4970	17329			
QKILIKIPVTKNIIT 2124 993 303 534 2240			713		536			5498	141			
VCLVICOVENEDEV 629 16 46393 17850	Y QKILIKIPVTKNIIT		993		303			534	2240			
	KCLVISQVSNSDSY		628		16			46383	17859			

			HL	A-DR SUPE	ERTYPE				
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901
K									
SKIMKLPKLPISNG	0400	824		6485			83674	110	
K FIHFFTWGTMFVPK	2126	745		273			489	1699	
Y LCNFKKNIIALLIIP	2127	9.7		312			423	21324	
KKNIIALLIIPPKIH	2128	13		203			495	157	
ALLIPPKIHISIEL	2129	648		1738			8.4	11957	
SMEYKKDFLITARK	2130	939		24			776	8897	
P	2131								
KSKFNILSSPLFNNF	2132	0.80		16			65	152	
FKKLKNHVLFLQM MN	2133	2.3		28			11	695	
KNHVLFLQMMNV	2100	12		32			757	>120098.04	
NLQ VLFLOMMNVNLQ	2134	6.3		30			8441	56770	
KQL	2135			30					
NVNLQKQLLTNHLI	0400	96		2460			555	11245	
N QKQLLTNHLINTPK	2136	675		228			4412	20984	
NHLINTPKIMPHHII	2137	1378		4798			625	1296	
YILLKKILSSRFNO	2138	220		183			8.3	18	
M	2139	400		2001			064	16604	
FNQMIFVSSIFISFY	2140	483		2091			854	16504	
KVSCKGSGYTFTA YQMH	2141	5000		381	50000		2946		
IAKVPPGPNITAEY		50000	>30000	>666.67	500000		>12500		
GDKWLD TAEYGDKWLDAKS	2142	50000	>30000	>666.67	16667		3125		
TWYGKPT AKSTWYGKPTGAG	2143	50000	>30000	667	500000		>12500		
PKDNGGA GAGPKDNGGACGY	2144	50000	>30000	>666.67	500000		>12500		
KDVDKAP	2145	50000	61062	× / / / / / /	500000		>12500		
FNGMTGCGNTPIFK DGRGCG	2146	50000	51962	>666.67	500000		>12500		
PIFKDGRGCGSCFEI		50000	6784	>666.67	500000		>12500		
KCTKP SCFEIKCTKPESCSG	<u>2147</u>	50000	>900000	>666.67	500000		12500		
EAVTV	2148								
AFGSMAKKGEEQN VRSAGEL	2149	50000	>30000	>666.67	50000		>12500		
TPDKLTGPFTVRYT		50000	>900000	>666.67	500000		>12500		
TEGGTK VRYTTEGGTKSEV	<u>2150</u>	50000	>30000	>666.67	500000		>12500		
EDVIPEG	2151								
TCVLGKLSQELHK LO	2152	26	29529	14848	7566	9001	18653	7656	17895
KLSQELHKLQTYPR	2153	19	196889	19684	2076	12198	85464	28656	19129
LHKLQTYPRTNTGS G	2154	2118	>205479.45	15182	9921	>7403.08	40226	1618	>29228.37
KLQTYPRTNTGSGT		>10060.36	>205479.45		114672	>9806.45	>99206.35	>51578.95	>29228.37
P CCVLGKLSQELHK	2155	34	17387	7 19 7 64	31253	5299	41656	5640	21704
LQ CSNLSTCVLGKLSQ	<u>2156</u>	296	>205479.45	14339	28603	5340	31837	3516	7225
E	2157								
TSNLSTTVLGKLSQ E	2158	298	86798	8016	32358	9280	31275	2058	2469
TTVLGKLSQELHKL	2158	133	92782	22449	36802	>9806.45	26113	16182	23824
Q DIAAKYKELGY	2159		>900000	>470.59					
ALVRQGLAKVA	2160 2161	1250		190	500000				
-	2101								

HLA-DR SUPERTYPE										
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901	
PATLIKAIDGDTVK		278	6429	296	3846		8333			
LMYKGQ TPETKHPKKGVEK	2162	>1000	>900000	>500	500000		12500			
YGPEASA VEKYGPEASAFTK	2163 2164	50000	>900000	1333	500000		12500			
KMVENAK FTKKMVENAKKIE VEFDKGQ	2165	>1000	11619	>500	500000		8333			
YIYADGKMVNEAL VROGLAK	2166	65		500	4167		1563			
HEQHLRKSEAQAK KEKLNIW	2167	50000	90000	80000	16667		6250			
QAKKEKLNIWSED NADSGQ	2168	50000	>900000	364	3125		>250000			
YFNNFTVSFWLRV PK	2169	50000		615	25000					
FSYFPS1	2170	50000		889	16667					
YSFFPSI	2171	50000		889	500000					
YSYFPSIR	2172	50000	>900000	667	16667		7217			
DPNANPNV DPNAN PNVNANPNANPNA				738	>5494.51		>15625			
NP(X4) QKWAAVVVPS	2173	50000		1000	50000					
TWQLNGEELIQDM	<u>2174</u>	50000		889	2273					
ELVETRPAG PEFLEQRRAAVDT	<u>2175</u>	5000		80000	500000		250000			
YC STORKUSP33	2176			617	2069					
DYSYLQDSDPDSFQ		>50000		189	>500000	>126666.67	>250000	>61250	>107142.8	
D DFSYLQDSDPDSFQ D	2178 2179			264	>500000	>126666.67	>250000	>61250	6 >107142.8 6	
QNILFSNAPLGPQF	2179			195					0	
QNILLSNAPLVPQF				538						
DYSYLQDSDPDSFQ	2181 2182			316	>166666.67					
KYVKQNTLKLAT	2183	9.9		6.2	25000					
P(X)KQNTLKLAT	2184	1.7							*****	
EEDIEIIPIQEEEY	2185	>9057.97	>18549.05	>7518.8	12203	849	>6742.18		128305	
HQAISPRTLNSPAIF YTDVFSLDPTFTIET	2186	1961	298315 217	6214	1314	3450	39701	14848	286179	
T YAGIRRDGLLLRLV	2187		9.6							
D LFFYRKSVWSKLQ	2188	19	30163	913	1383	84	84	65		
SI RPIVNMDYVVGAR	2189	29	22	3.1	21	812	346	748		
TFRREKR RPGLLGASVLGLD	2190	1789	35768	6522	4414	3183	506	>61250		
DI LYFVKVDVTGAYD	<u>2191</u>	16	9.6	2.8	13	14	5892	413		
TI FAGIRRDGLLLRLV	2192	2381	3.6	7092	3820	>3365.21	41148	7650		
D AKTFLRTLVRGVPE	2193	104	54159	208	3326	105	25	9.2		
Y YGAVVNLRKTVVN	2194	13509	150175	4194	4531	>95000	8274	113		
FP GTAFVQMPAHGLF PW	2195 2196	1.6	37275	8.1	34	18	90	99		
WAGLLLDTRTLEV		2016	22	49	323	1238	186	>61250		
QS RTSIRASLTFNRGF K	2197 2198	1430	256	770	177	5131	411	5475		

	HLA-DR SUPERTYPE												
Sequence	SEQ ID	DRB1 *0101	DRB1 *0301	DRB1 *0401	DRB1 *0404	DRB1 *0405	DRB1 *0701	DRB1 *0802	DRB1 *0901				
RVIKNSIRLTL	2199	3650	584	9249	5389	80682	2239	1175	2566				
PVIKNSIKLRL	2200	1549	198	34245	14612	277735	4091	541	2851				
ATSTKKLHKEPATL IKAIDG		4.6	8018	113	1020		2083						

TABLE 27

	CEO		A-DR SUPI		DDD2	DDB4	DDD#	DRB
Sequence	SEQ ID NO.	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	*0201
AC- NPTKHKWEAAHVAEQ		>33333.33	>10000	200000	101		1250	
LAA	<u>1543</u>	×1111 11			>11111.11		35	
DDYVKQYTKQYTKQN TLKK	1544	>1111.11			>111111.11		33	
AAAKAAAAAAYAA	1545	200000			6250		2857	
AC-								
AAAKAAAAAAYAA (20)AYA(20)A(20)A(20)K (20)A(20)	1546 1547	200000					2857	
AC-AAAKATAAAAYAA	1548							
ΑС-ΑΑΑΚΑΑΑΑΑΑΑΓΑΑ								
AC- AAAKATAAAA(10)AA	1549 1550							
AC-								
AAAKATAAAA(23)AA AAKAAAAAAA(10)AA	1551							
AAYAAAATAKAAA	1552							
AALAAAAAAKAAA	1553	2222					67	
AAEAAAATAKAAA	<u>1554</u>	2222					0,	
	1555							
AAYJJAAAAKAAA	<u>1556</u>							
AAYAAAAJJKAAA	1557							
AFLRAAAAAAFAA	1558							
AFLRQAAAAAFAAY	1559							
AAFAAAKTAAAFA	1560	4.6			20000		25	6.4
YAAFAAAKTAAAFA	1561	2.6			33333	30	9.5	
AALKATAAAAAAA	1562							
YAR(15)ASQTTLKAKT	1563	3.9					3.6	
YARF(33)QTTLKAKT	1564							
PKYFKQRILKFAT	1565							
PKYFKQGFLKGAT	1566							
PKYGKQIDLKGAT								
AAFFFFFGGGGGA	1567							
AADFFFFFFFDA	1568							
AAKGIKIGFGIFA	1569							
AAFIFIGGGKIKA	1570							
AAKIFIGFFIDGA	1571							
AAFIGFGKIKFIA	1572							
	1573							
AAKIGFGIKIGFA	1574							
AAFKIGKFGIFFA	1575							
AADDDDDDDDDA	1576							
(43)AAIGFFFFKKGIA	1577							
(43)AAFFGIFKIGKFA	1578							
(43)AADFGIFIDFIIA	1579							
(43)AAIGGIFIFKKDA	1580							
(43)AAFIGFGKIKFIA	1581							
(43)AAKIGFGIKIGFA	1582							
(43)AAFKIGKFGIFFA	1583							

	SEQ	DRB1	LA-DR SUPE DRB1	DRB1	DRB3	DRB4	DRB5	DRB5
Sequence	ID NO.	*1101	*1302	*1501	*0101	*0101	*0101	*0201
AAAKAAAAAAAFA	1585							
AAAKAAAAAAFAA	1586							
AAAKAAAAFAAAA	1587							
FAAAAAAAAAAA	1588							
AAAAAAAAAAA	1589							
AAAAAAAAAANA	1590							
AAANAAAAAAAA	1591							
AAAAAAAAAAA	1592							
AAAAASAAAAAA	1593							
ASAAAAAAAAA	1594							
AFAAAKTAA								
YARFLALTTLRARA	1595							
YAR(15A)SQTTLKAKT	1596	2.5					1.4	48
YAR(15A)RQTTLKAAA	1597	1.2					0.94	62
(15A)RQTTLKAAA	1598	1.8					9.5	3095
(16A)RQTTLKAAA	1599	77					4000	
	1600	"					-1000	
(46)AAKTAAAFA	1601							
(39)AAAATKAAA	1602							
(52)AAAATKAAAA	1603							
(55)AAAATKAAAA	1604						120	
A(14)AAAKTAAA	1605	43					120	
AA(14)A(35)ATKAAAA	1606							
AA(14)AA(36)TKAAAA	1607							
AFAAAKTAA(72)	1608							
(49)AAAKT(64)AAA	1609							
(49)AAAKTA(64)AA	1610							
HQAISPRTLNGPGPGSP		9875	638	5570		232	32930	
AIF YAAFAAAKTAAAFA	1611				>4347.83			
TEGRCLHYTVDKSKPK	1612	>1250			4082		2857	
AWVAWRNRCK	1613	>5000			>11111.11		44	
IVSDGNGMNAWVAWR	1614	6667			>6250		>2222.22	
NRC	1615							
PHHTALRQAILSWGEL MTI.A	1616	3116		5.3		48	261	
WMYYHGQRHSDEHHH	1617	>10000			>7692.31		>5000	
YIVMSDWTGGA		>6666.67			>33333.33		>10000	
АНААНААНААНА	<u>1618</u>	200000					200000	
A MDIDPYKEFGATVELLS	<u>1619</u>		2415					
MDIDPYKEFGATVELLS FLPSDFFP	1620		2415					
GMLPVCPLIPGSSTTST		2500			>25000		200000	
GP LGFFPDHQLDPAFRANT	1621	6667			1449		6667	
GYKVLVLNPSV	1022	26	21	126		995	>11441.65	
LMAFTAAVTS	1623	>23337.22	>2464.79	1934		11687	>12586.53	
TFALWRVSAEEY	1624	342	>2569.75	>12709.5		>6608.93	25499	
ALWRVSAEEY	1625	243	>6398.54	>15268.4		>7930	>35587.19	
	1626			6				
EEYVEIRQVGDFH	1627	4683	>1895.99	2060		2063	9754	
VGGVYLLPRRGPRLGV	1628	88	>15350.88	4.2	60753	19239	12	

	SEQ	DRB1	LA-DR SUPE DRB1	DRB1	DRB3	DRB4	DRB5	DRB
Sequence	ID NO.	*1101	*1302	*1501	*0101	*0101	*0101	*020
VGGAYLLPRRGPRLGV	1629	507	24663	4.1	>66533.6	37640	50	
VGGVALLPRRGPRLGV	1630	154	>15350.88	8.5	>66533.6	25688	20459	
VGGVYALPRRGPRLGV	1631	12	>15350.88	451	>66533.6	26122	34	
VGGVYLAPRRGPRLGV	1632	35	>15350.88	55	>66533.6	>42059.4 6	76	
VGGVYLLARRGPRLGV	1633	6.5	10325	2.8	17030	4338	17	
VGGVYLLPARGPRLGV	1634	694	201	6.5	18073	18960	40	
VGGVYLLRRAGPRLGV	1635	67	>15350.88	6.2	91912	30707	7.9	
GAPLGGAARALAHGV	1636	24	8739	1615	>70972.32	3959	11983	
GAALGGAARALAHGV	1637	168	19335	4483	>70972.32	3509	25372	
GAPLAGAARALAHGV	1638	9.5	7215	2810	>70972.32	2963	7688	
GAPLGAAARALAHGV	1639	36	15091	3920	>70972.32	16533	4502	
GAPLGGLARALAHGV		12	76	1805	123762	3950	4256	
GAPLGGALRALAHGV	1640	83	340	2068	>51098.62	4889	5396	
GAPLGGAAAALAHGV	1641	43842	23810	7682	>51098.62	31	12916	
GAPLGGAARLLAHGV	1642	80	29412	631	>51098.62	2549	26684	
GAPLGGAARAAAHGV	1643	3633	>23489.93	>8666.67	>51098.62	41441	42463	
GAPLGGAARALAAGV	1644	45	23179	5714	>51098.62	3865	8354	
FPDWQNYTPGPGTRF	1645	>51282.05	>12027.49	35058		33923	>20533.88	
RFPLTFGWCFKLVPV	1646	62289	4797	514		964	>20533.88	
RQDILDLWVYHTQGY	1647	>51282.05	6775	723		1326	16155	
RQEILDLWVYHTQGF	1648	11113	5384	985		1071	>20533.88	
LSHFLKEKGGLEGLI	1649	9460	>12027.49			18709	>20533.88	
	1650			9				
LSFFLKEKGGLDGLI	1651	614	>12027.49	>39737.9		13214	15272	
LEPWNHPGSQPKTACT	1652	>15325.67	>11041.01	2665		92	2939	
QVCFITKGLGISYGR	1653	31	92	3555		876	3950	
QLCFLKKGLGISYGR	1654	9.5	88	4212		282	1190	
PPEESFRFGEEKTTPS	1655	>10000			>14285.71		>2857.14	
CIVYRDGNPYAVCDK	1000	>14662.76	1646	650		>24786.3	>10666.67	
HYCYSLYGTTLEQQY	1656	12397	>13725.49	4849		2 1292	>10666.67	
	1657	>14662.76	>13725.49	5060		189	>10666.67	
CYSLYGTTLEQQYNK	1658							
NTSLQDIEITCVYCK VFEFAFKDLFVVYRD	1659	>14662.76 10923	14857 7675	678 4871		11710 18117	>10666.67 >10666.67	
	1660							
EFAFKDLFVVYRDSI	1661	9496	9996	5355		9072	5998	
DLFVVYRDSIPHAAC	1662	1163	11172	2832		2676	10741	
FVVYRDSIPHAACHK	1663	1194	1851	349		18144	2343	
NTGLYNLLIRCLRCQ	1664	14	5692	67		222	598	
IRCLRCQKPLNPAEK	1665	>14662.76	>13725.49	6928		611	>10666.67	
PRKLHELSSALEIPY	1666	5990	51	1116		1710	>10666.67	
EIPYDELRLNCVYCK	1667	>18001.8	858	2084		9047	>62305.3	
TEVLDFAFTDLTIVY	1668	>18001.8	>13059.7	561		110	>62305.3	
VLDFAFTDLTIVYRD	1669	7474	3102	645		11294	14839	
DFAFTDLTIVYRDDT	1670	14334	5008	3651		21621	675	
TIVYRDDTPHGVCTK		>18001.8	6280	5449		>21521.3 4	>62305.3	
WYRYSVYGTTLEKLT	1671	1670	805	421		1039	62	
	1672	>18001.8				112	>62305.3	

_	SEQ	DRB1	DRB1	DRB1	DRB3	DRB4	DRB5	DRB
Sequence	ID NO.	*1101 >18001.8	*1302 >13059.7	*1501 955	*0101	*0101 1325	*0101 11802	*020
SEVYDFAFADLTVVY	1674			933		10720	27275	
VYDFAFADLTVVYRE	1675	>18001.8	>13059.7				39785	
DFAFADLTVVYREGN	1676	>18001.8	9627	4915		17973		
TVVYREGNPFGICKL	1677	>18001.8	>13059.7	13850		16200	48840	
GNPFGICKLCLRFLS	1678	1084	9737	1139		196	6594	
NYSVYGNTLEQTVKK	1679	>56657.22	8614	15587		>25108.2	14326	
KKPLNEILIRCIICQ	1680	1299	965	1870		140	26273	
NEILIRCIICQRPLC	1681	20827	7174	18927		883	>29761.9	
IRCIICQRPLCPQEK		6757	7295	25349		510	15154	
CIVYRDCIAYAACHK	1682	35566	12898	3847		2578	1912	
NTELYNLLIRCLRCQ	1683	259	5674	2449		797	854	
IRCLRCOKPLNPAEK	1684	21581	>9641.87	27591		447	20171	
REVYKFLFTDLRIVY	<u>1685</u>	2263	80	258		203	155	
RIVYRDNNPYGVCIM	1686	3446	119	821		1403	20474	
NNPYGVCIMCLRFLS	1687	7786	4797	6662		207	7258	
EERVKKPLSEITIRC	1688	6877	8919	132		2990	7910	
IRCIICQTPLCPEEK	1689	5461	17444	9766		916	>51020.41	
-	1690						16853	
EIPLIDLRLSCVYCK	<u>1691</u>	47355	6936	656		861		
SCVYCKKELTRAEVY	1692	569	23385	4374		673	3197	
VCLLFYSKVRKYRYY	1693	326	309	61		2343	182	
YYDYSVYGATLESIT	1694	9122	8923	1106		32378	>51020.41	
IRCYRCQSPLTPEEK	1695	6645	>14403.29	480		28659	>51020.41	
VYDFVFADLRIVYRD	1696	12168	79	855		4392	>51020.41	
DFVFADLRIVYRDGN	1697	6957	162	1253		6709	8433	
RIVYRDGNPFAVCKV	1698	174	122	81		1606	3148	
GNPFAVCKVCLRLLS	1699	296	7389	117		126	657	
KKCLNEILIRCIICQ	1700	7579	731	3176		257	>9925.56	
NEILIRCIICQRPLC	1701	16056	10184	8177		372	>22909.51	
RTAMFQDPQERPRKL	1702	1034	17086	73192		20481	7474	
LFVVYRDSIPHAACH	1703	1582	697	437		3580	7854	
LTIVYRDDTPHGVCT	1704	15880	1852	27048		16993	>15267.18	
LCIVYRDCIAYAACH	1705	9886	5662	2269		2881	9738	
YKFLFTDLRIVYRDN	1706	10122	77	2912		1342	800	
YNFACTELKLVYRDD		11615	10167	3082		12866	1673	
LKLVYRDDFPYAVCR	1707	698	699	1877		3828	9156	
YDFVFADLRIVYRDG	1708	6540	8173	25727		10907	11161	
LRIVYRDGNPFAVCK	1709	109	123	169		1566	6820	
HEYMLDLQPETTDLY	1710	>56179.78	12990	30895		2099	>22909.51	
TLRLCVOSTHVDIRT	1711	17613	932	3957		243	>22909.51	
IRTLEDLLMGTLGIV	1712	1156	789	2181		23	12385	
LEDLLMGTLGIVCPI	1713	8514	1693	229		1800	9475	
DLLMGTLGIVCPICS	<u>1714</u>	>56179.78	1053	1427		4123	16198	
	<u>1715</u>	25948	603	6968		159	>9925.56	
KATLQDIVLHLEPQN	<u>1716</u>							
IDGVNHQHLPARRAE	1717	>56179.78	>11475.41	1		344	12573	
LRAFQQLFLNTLSFV	1718	106	1.01	20		2.2	253	
FQQLFLNTLSFVCPW		10311	9.3	24792		309	17330	

	SEQ	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	DRB5 *0201
Sequence ODYVLDLQPEATDLH	ID NO.	>11918.95	*1302 >11475.41	>62758.6	*0101	1851	>22909.51	-0201
•	1720			2				
DIRILQELLMGSFGI	1721	18982	5796	1625		16	>55096.42	
RILQELLMGSFGIV	1722	7978	1038	294		17	>55096.42	
ELLMGSFGIVCPNCS	1723	>59171.6	933	1928		206	>55096.42	
KEYVLDLYPEPTDLY	1724	>59171.6	>14767.93	3171		476	>55096.42	
LRTIQQLLMGTVNIV	1725	3641	6.4	265		15	32108	
QQLLMGTVNIVCPT	1726	11062	9.0	2010		166	>55096.42	
QLLMGTVNIVCPTCA	1727	>59171.6	118	>38396.6		11550	>55096.42	
RETLQEIVLHLEPQN	1727	7896	11360	16220		95	>55096.42	
LRTLQQLFLSTLSFV	1728	208	55	29		3.1	1994	
LQQLFLSTLSFVCPW	1729	11693	133	296		22	36943	
KDYILDLQPETTDLH	<u>1730</u>	>17436.79	23654	>37448.5		490	>55096.42	
-	<u>1731</u>			6				
LRTLQQMLLGTLQVV	1732	907	616	1697		88	>46620.05	
LQQMLLGTLQVVCPG	1733	>31645.57	395	1266		1014	29198	
QMLLGTLQVVCPGCA	1734	>31645.57	874	4144		258	>31446.54	
VPTLQDVVLELTPQT	1735	>31645.57	14985	12263		1000	>31446.54	
LQDVVLELTPQTEID	1736	>31645.57	1145	>33090.9		1116	>31446.54	
QDVVLELTPQTEIDL		>31645.57	10274	>33090.9		1719	>31446.54	
CKFVVQLDIQSTKED	1737	>31645.57	>11437.91	1 22851		301	>31446.54	
VVQLDIQSTKEDLRV	<u>1738</u>	7353	708	5044		226	8690	
DLRVVQQLLMGALTV	1739	667	57	132		9.5	10879	
LRVVQQLLMGALTVT	1740	314	8.9	56		7.7	8755	
VOOLLMGALTVTCPL	1741	11074	574	526		204	7151	
	1742	7657	1223	4461		1470	>31446.54	
QQLLMGALTVTCPLC	1743							
QLLMGALTVTCPLCA	1744	>31645.57	1817	3761		2224	>31446.54	
REYILDLHPEPTDLF	1745	4152	13183	>33090.9		316	>31446.54	
CCYTCGTTVRLCIN	1746	8636	739	3820		891	16033	
VRTLQQLLMGTCTIV	1747	1409	37	1829		139	>15267.18	
LQQLLMGTCTIVCPS	1748	9447	753	2441		2667	>15267.18	
MLDLQPETTDLYCYE		>15209.13	>12027.49			20	>15267.18	
VLDLYPEPTDLYCYE	1749	>15209.13	>12027.49	6 21591		18	>15267.18	
LREYILDLHPEPTDL	1750	9827	12365	10949		2040	>40404.04	
HIEFTPTRTDTYACRV	1751	200000	12303	10,47	>7142.86	2040	200000	
LWWVNNESLPVSPRL	1752	200000			- 77 12.00		200000	
YEEYVRFDSDVGE	1753	200000					200000	
EEYVRFDSDVGE	1754	200000					200000	
APPRLICDSRVLERY	1755	>1111111.11	149	1384	1617	2840	6087	
	1756	2945	20402	85	16159	8550	7295	
CDSRVLERYLLEAK	<u>1757</u>		881		340			
VLERYLLEAKEAENI	<u>1758</u>	17227		269	340 8307	8920 52943	6714 6626	
EHCSLNENITVPDTK	1759	>	84	12013				
NENITVPDTKVNFYA	1760	17921	9338	22568	>38167.94	>38461.5 4	12214	
VPDTKVNFYAWKRME	1761	8861	14795	333	>38167.94	23602	449	
VNFYAWKRMEVGQQA	1762	50	14798	1194	22507	1490	455	
WKRMEVGQQAVEVW	22.00	512	159	1812	>42194.09	238	4300	

			A-DR SUPE				none.	nnn-
Sequence	SEQ ID NO.	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	DRB5 *0201
VGQQAVEVWQGLALL	1764	>17241.38	1313	12	>38167.94	3901	>7785.13	
VEVWQGLALLSEAVL	1765	5157	4473	58	>38167.94	1334	13794	
GLALLSEAVLRGQAL	1766	2578	1216	1939	>38167.94	3.5	105	
SEAVLRGQALLVNSS	1767	3484	7.4	151	3997	23	1057	
RGQALLVNSSQPWEP	1768	7698	3.4	2876	6165	1554	558	
LVNSSQPWEPLQLHV	1769	>8163.27	504	2359	18044	3412	10039	
QPWEPLQLHVDKAVS	1770	8897	695	12480	1924	103	2929	
LQLHVDKAVSGLRSL	1771	910	53	2707	1044	31	76	
DKAVSGLRSLTTLLR	1772	52	187	60	3150	2006	104	
GLRSLTTLLRALGAQ	1773	3.7	871	6.2	12947	283	2.7	
TTLLRALGAQKEAIS	1774	860	1512	89	33256	251	21	
ALGAQKEAISPPDAA	1775	4212	>12411.35	14216	>91743.12	27294	3963	
KEAISPPDAASAAPL	1776	601	9272	1201	27203	2988	310	
PPDAASAAPLRTITA	1777	2582	10205	1267	10584	182	1117	
SAAPLRTITADTFRK	1778	3883	809	858	2111	17	45	
RTITADTFRKLFRVY	1779	166	95	35	672	1561	93	
DTFRKLFRVYSNFLR	1780	11	10	0.95	43687	1029	26	
LFRVYSNFLRGKLKL	1781	173	80	2.8	8981	2333	2.9	
SNFLRGKLKLYTGEA	1782	192	4730	30	4075	2442	5.7	
KLKLYTGEACRTGDR	1783	>17241.38	880	130	17787	20089	636	
APPRLITDSRVLERY	1784	2750	92	238	710	2263	698	
ITDSRVLERYLLEAK	1785	5279	>14705.88	18	>42194.09	12401	621	
EHTSLNENITVPDTK		>408163.27	13	11082	>42194.09	>29029.0	5547	
KLKLYTGEATRTGDR	1786 1787	4364	841	18	5298	3 14838	731	
PQPFRPQQPYPQ	1788					15		
PFRPQQPYPQ	1789					42		
PQPFRPQQPYP	1790					14		
PQPFRPQQP	1791					19		
KQPFRPQQPYPQ	1792					56		
PKPFRPQQPYPQ	1793					3.4		
PQPFKPQQPYPQ	1794					19		
PQPFRKQQPYPQ	1795					22		
PQPFRPQKPYPQ	1796					22		
PQPFRPQQPKPQ	1797					325		
PQPFRPQQPYKQ	1798					35		
PQPFRPQQPYPK	1799					22		
QFLGQQQPFPPQ	1800					2.8		
FLGQQQPFPPQ	1801					31		
LGQQQPFPPQ	1802					151		
QFLGQQQPFPP	1803					2.3		
QFLGQQQPF	1804					5.3		
IRNLALQTLPAMCNVY	1805					1.9		
NLALQTLPAMCNVY						27		
NEADQTEI AMCHTT								
LALQTLPAMCNVY	1806 1807					153		
						153 2.0 3.0		

			LA-DR SUPI					
Sequence	SEQ ID NO.	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	DRB5 *0201
EGDAFELTVSCQGGLP		1101	1502	1501	0.01	0.0.		0201
K ESTGMTPEKVPVSEVM	1810		>17500			>64444.4		
GT	1811					4		
FPTIPLSRLFDNASL	1812	30675	7495	1390	2585	194	5799	
RLFDNASLRAHRLHQ	1813	12461	84	85	11411	3210	557	
LRAHRLHQLAFDTYQ	1814	3208	7590	90	19811	2.0	4471	
QLAFDTYQEFEEAYI	1815	>15384.62	15167	23166	595	11495	>38610.04	
QEFEEAYIPKEQKYS	1816	12821	>15837.1	>15582.1	>54554.47	>41134.7	5418	
IPKEQKYSFLQNPQT	1817	>15384.62	13695	16207	30572	55587	13118	
SFLQNPQTSLCFSES	1818	>15384.62	190	6513	93809	21651	>9647.76	
TSLCFSESIPTPSNR	1819	>15384.62	99	1944	3920	1883	>38610.04	
REETQQKSNLELLRI	1820	>15384.62	15709	9736	>270270.27	52	25133	
SNLELLRISLLLIQS	1821	23669	196	59	>91901.83	147	50110	
ISLLLIQSWLEPVQF	1822	2675	120	60	6765	2.5	>9960.16	
SWLEPVQFLRSVFAN	1823	2715	4322	136	>270270.27	291	4815	
FLRSVFANSLVYGAS	1824	973	5.6	13	157978	814	141	
NSLVYGASDSNVYDL	1825	>15384.62	14038	3640	11769	1792	>13046.31	
SDSNVYDLLKDLEEG		>15384.62	>17857.14	>30536.9	219298		>13046.31	
GIQTLMGRLEDGSPR	1826	4474	10433	1 1348	186220	22 2110	18006	
RLEDGSPRTGQIFKQ	1827	7896	>17857.14	9106	18119	296	12580	
RTGQIFKQTYSKFDT	1828	6961	66	155	14736	201	64	
OTYSKFDTNSHNDDA	1829	>15384.62	>17857.14	25883	38715	>137767.	5787	
TNSHNDDALLKNYGL	1830 1831	>15384.62	5169	133	130378	22 >137767. 22	>13046.31	
ALLKNYGLLYCFRKD	1832	>15384.62	10	17	2309	1230	462	
DMDKVETFLRIVQCR	1833	885	1232	201	>27322.4	826	7447	
FLRIVQCRSVEGSCGF	1834	2708	1017	839	>27322.4	1078	7102	
FPTIPLSRLFDNAML	1835	46404	9313	2770	121212	216	11521	
RLFDNAMLRAHRLHQ	1836	267	738	18	>270270.27	1628	58	
QLAFDTYQEFEQNPQ		>15384.62	19718	>86666.6	738	>32842.5	>9510.22	
SFLQNPQTSLCCFRK	1837	3801	128	7 103	>270270.27	8 8500	3739	
SNLELLRICLLLIQS	1838	>15384.62	773	90	17024	164	>11771.33	
ICLLLIQSWLEPVQF	1839	>15384.62	954	1771	187970	49	>9510.22	
NSLVYGASDSNIYDL	1840	>15384.62	10854	971	31616	3287	>9510.22	
SDSNIYDLLKDLEEG	1841	>15384.62	>16203.7	>86666.6	>18726.59	24259	>9510.22	
	1842			7				
DKVETFLRIVQCCGF	1843	1023	1034	383	6278	184	6350	
SFLQNPQTSLTFSES	1844	>15384.62	121	1511	864	17824	12365	
TSLTFSESIPTPSNR	1845	22152	16	176	>95238.1	3476	>1335.38	
ALLKNYGLLYTFRKD	1846	1737	0.89	6.5	50	1335	29	
LLYTFRKDMDKVETF	1847	7905	>14522.82	886	941	12493	154	
DMDKVETFLRIVQTR	1848	206	3381	>86666.6 7	13712	190	1263	
FLRIVQTRSVEGSTGF	1849	143	1.5	9.8	27345	21	116	
HLDMLRHLYQGCQVV	1850	2076	2879	359	107066	163	7087	
RLRIVRGTQLFEDNYAL	1851	2072	5.2	31	1198	120	46	
GVGSPYVSRLLGICL	1852	696	955	46	148588	316	14197	
TLERPKTLSPGKNGV	1853	>52631.58	835	23264	>263157.89	25739	11337	
	1000							

	HLA-DR SUPERTYPE										
Seguence	SEQ ID NO.	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	DRB5 *0201			
Sequence KIFGSLAFLPESFDGDP	ID NO.	>52631.58	1073	2264	43745	10020	8008	0201			
A	1854	4573	>71428.57	7891	15838	970	4055				
ELVSEFSRMARDPQ	1855	4573 79	29	269	13636	1023	46				
GEALSTLVLNRLKVG	1856	79 51	29 816	489		902	4517				
AYVLLSEKKISSIQS VASLLTTAEVVVTEI	1857	>18674.14	>10294.12	>50837.9			>119047.6				
VASLLTTAEVVVTEI	1858	>180/4.14	>10294.12	9		3	2				
KCEFQDAYVILLSEKK	1859	1078	>10294.12	>47643.9 8		>19594.5	20				
ALSTLVLNRLKVGLQ	1860	9.1	4.6	191		17	3.9				
MSYNLLGFLQRSSNC	1861	3628	1190	89	>42194.09	6503	710				
LGFLQRSSNCQCQKL	1862	6025	112	1397	>42194.09	1167	649				
RSSNCQCQKLLWQLN	1863	>408163.27	6153	802	3519	21	6981				
QCQKLLWQLNGRLEY	1864	1644	227	175	8709	209	924				
LWQLNGRLEYCLKDR	1865	4215	808	893	29028	15576	3241				
GRLEYCLKDRRNFDI		1707	1240	940	5213	15870	64725				
RNFDIPEEIKQLQQF	1866	7326	>15418.5	2036	23832	311	6854				
PEEIKQLQQFQKEDA	1867	1953	13325	1873	>26315.79	215	675				
OLOOFOKEDAAVTIY	1868	>408163.27	68	1724	348	1338	4270				
OKEDAAVTIYEMLON	1869	>408163.27	7315	1146	>42194.09	15173	>10482.18				
AVTIYEMLQNIFAIF	1870	29718	109	262	2828	1118	14047				
EMLQNIFAIFRQDSS	1871	36832	61	1718	726	164	3187				
IFAIFRODSSSTGWN	1872	4558	775	204	2181	30	109290				
RODSSSTGWNETIVE	1873	>42553.19	848	>189583.	9172	1497	8650				
•	1874	20576	105	33 897	>26315.79	166	5822				
STGWNETIVENLLAN	1875	20576	8.5	1603	>42194.09	2503	18559				
ETIVENLLANVYHQR	1876	>42553.19 8258	8.5 61	20	>123456.79	3071	65				
NLLANVYHQRNHLKT	1877	22002	1267	1662	>123456.79	9585	4.7				
VYHQRNHLKTVLEEK LEKEDFTRGKRMSSL	1878	698	25362	14118	6267	16057	4903				
FTRGKRMSSLHLKRY	1879	81	10245	118	18836	2027	84				
RMSSLHLKRYYGRIL	1880	1035	2532	1.3	>26178.01	2255	491				
	1881				6608	2233					
HLKRYYGRILHYLKA	1882	2721	868	0.69			2.3				
YGRILHYLKAKEDSH	1883	812	2783	16	454545	140	39				
HYLKAKEDSHCAWTI	1884	>60606.06	11571	627	301205	7501	2632				
KEDSHCAWTIVRVEI	1885	9320	506	1397	>1754385.9 6	7.9	4056				
CAWTIVRVEILRNFY	1886	4167	147	196	10300	152	4143				
VRVEILRNFYVINRL	1887	504	5.8	1.04	80386	187	485				
RNFYVINRLTGYLRN	1888	55	9.4	18	689	1249	5.6				
MSYNLLGFLQRSSNT	1889	3069	1334	6.8	51787	4660	9.0				
LGFLQRSSNTQTQKL		26247	21	2331	>1754385.9	1041	339				
RSSNTQTQKLLWQLN	1890	>42553.19	169	2740	6 751	26	8545				
QTQKLLWQLNGRLEY	1891	20654	121	20	6582	88	417				
LWQLNGRLEYTLKDR	1892	6521	2447	853	4402	14310	6004				
GRLEYTLKDRRNFDI	1893	4998	1468	168	9901	21427	796				
HYLKAKEDSHTAWTI	1894	>60606.06	2264	529	35829	11750	19617				
KEDSHTAWTIVRVEI	1895	7443	3046	1992	56205	18	575				
TAWTIVRVEILRNFY	1896 1897	5052	72	242	14419	26	518				
LGFLQRSSNCQSQKL	1897	604	131	541	>1754385.9	124	508				
	1070										

	SEQ	DRB1	DRB1	DRB1	DRB3	DRB4	DRB5	DRB
Sequence	ID NO.	*1101	*1302	*1501	*0101	*0101	*0101	*020
RSSNCQSQKLLWQLN		>60606.06	1960	2962	68823	27	4077	
OSOKLLWOLNGRLEY	1899	>60606.06	155	108	5609	166	402	
GIVEOCCTSICSLYO	1900	7940	239	1280	14353	4245	>37593.98	
TSICSLYQLENYCN	1901	>10526.32	>15021.46	837	8048	13496	>40322.58	
GILEOCCTSICSLYQ	1902	>10526.32	858	1097	>18726.59	5871	19231	
GIVEQTTTSITSLYQ	1903	>10526.32	14	849	>95238.1	2303	>37593.98	
EOTTTSITSLYQLEN	1904	>10526.32	16949	1078	>18726.59	29614	48505	
	1905	>10526.32	10346	173	>95238.1	1645	>40322.58	
TSICSLYQLENYCG	1906	1095	>17073.17	99	>95238.1	3245	6048	
TSITSLYQLENYTN	1907	1093	>17073.17	182	92336	1658	16073	
TSITSLYQLENYTG	1908		15347	237	14184	11017	>43290.04	
GIVEQCCCGSHLVEA	1909	>10526.32						
SLYQLENYCCGERGF	1910	>11111111.11	>15909.09	151	92336	30978	>43290.04	
CCTSICSLYQLENYCC	<u>1911</u>	>11111111.11	7096	877	>18726.59	1582	>40650.41	
GSHLVEALYLVCCN	1912	>1111111.11	3259	11191	>18726.59	14065	>46403.71	
CCGSHLVEALYLVCC	1913	>10526.32	6027	12986	>18726.59	11357	>43290.04	
FVNQHLCGSHLVEAL	1914	>1111111.11	10595	1195	>95238.1	3153	47170	
QHLCGSHLVEALYLV	1915	>10526.32	7624	103	14819	1480	32049	
GSHLVEALYLVCGER	1916	>10526.32	8030	1350	>18726.59	372	29283	
VEALYLVCGERGFFY	1917	3563	4403	181	4443	30	25543	
YLVCGERGFFYTPKT	1918	>10526.32	9272	10655	92764	34450	95238	
FVNQHLCGSDLVEAL	1919	>11111111.11	20248	9679	10031	24511	>43290.04	
FVNQHLTGSHLVEAL	1920	>10526.32	12413	799	94518	4084	>43290.04	
QHLTGSHLVEALYLV	1921	>10526.32	6862	184	4027	939	23716	
GSHLVEALYLVTGER	1922	>10526.32	12185	1429	18215	225	11398	
VEALYLVCGERGSFY	1923	>10526.32	4288	1240	>95238.1	129	804	
VEALYLVCGERGFLY	1924	55402	1871	149	843	19	5149	
VEALYLVTGERGFFY	1925	4860	1076	116	17156	13	78	
YLVCGERGFLYTPKT	1926	>1111111.11	2120	>25633.8	>95238.1	33114	971	
YLVCGERGFFYTDKT		>60606.06	1014	>25633.8	616	48099	>28449.5	
YLVCGERGFFYTKPT	1927	>60606.06	3467	>25633.8	12805	40379	>28449.5	
YLVTGERGFFYTPKT	1928	7625	2100	>25633.8	13737	20721	>28449.5	
YLVTGERGFFYTDKT	1929	16849	17353	>25633.8	359	30824	>28449.5	
YLVTGERGFFYTKPT	1930	9341	17869	>21016.1	9573	27915	11926	
	1931			7				
VCGERGFFYTPKTRR	1932	3817	34669	>25633.8	17416	>30999.4 7	92	
VTGERGFFYTPKTRR		10116	25362	2824	243902	>29820.0	540	
MWDLVLSIALSVGCT	1933	81096	108	11375	15205	5 158	70711	
DLVLSIALSVGCTGA	1934	>200000	98	18200	>14918.69	459	>100000	
HPQWVLTAAHCLKKN	1935	981	483	1219	8114	1106	11	
QWVLTAAHCLKKNSQ	1936	14213	>35000	>45500	>14918.69	14395	382	
GORVPVSHSFPHPLY	1937	>200000	703	3960	>14918.69	9860	>200000	
RVPVSHSFPHPLYNM	1938	>200000	377	5518	>14918.69	9213	11650	
	1939	6455	3307	3873	>14918.69	49	1901	
PHPLYNMSLLKHQSL	1940			3873 472				
HPLYNMSLLKHQSLR	1941	248	546	>30333.3	>14918.69	8.4 105	219 >100000	
NMSLLKHQSLRPDED	1942	25820	>35000	>30333.3	~14916.09	105	-100000	

		HL	A-DR SUP	ERTYPE				
Sequence	SEQ ID NO.	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	DRB5 *0201
SHDLMLLRLSEPAKI		5267	1.8	365	5361	10	2031	0201
HDLMLLRLSEPAKIT	1943	1147	0.83	115	488	12	211	
PEEFLRPRSLQCVSL	1944	10675	11667	3193	>14413.38	117	57537	
PRSLQCVSLHLLSND	1945	11128	3731	1597	11650	544	46416	
NGVLQGITSWGPEPC	1946	32444	>17500	835	>14413.38	5761	>100000	
KPAVYTKVVHYRKWI	1947	327	1947	401	7186	4581	23	
LHLLSNDMCARAYSE	1948	26012	1876	>2367.33	1308	324	28817	
VGNWQYFFPVIFSKA	1949	20012	1070	- 2507.55			20011	
ESEFQAALSRKVAKL	1950							
IGHLYIFATCLGLSYDG	<u>1951</u>							
L	1952							
VGNWQYFFPVIFSKAS DSLQLVFGIELMEVD	1953							
PAYEKLSAEQSPPPY								
RNGYRALMDKSLHVG	1954							
TQCALTRR	1955							
FFKNIVTFFKNIVT	<u>1956</u>	****					1333	2065
YKSAHKGFKGVDAQG TLSKI	1957	2000					1333	2003
VDAQGTLSKIFKLGGR		18			769		6667	1152
DSRS AC-	1958	200000					200000	4561
ASQKRPSQRHGSKYLA								
TAST ENPVVHFFKNIVTPR	1959			5.2			463	
ENPVVAFFKNIVTPR	1960			2.8			302	
ENPVVHAFKNIVTPR	1961			4.1			910	
ENPVVHFFANIVTPR	1962			2.9			6235	
ENPVVHFFKNIVTPA	1963			2.5			3333	
NPVVHFFKNIVT	1964			23			10000	
HFFKNIVTPRTPPY	1965			460			377	
	<u>1966</u>			3.7			1890	
NPVVHFFKNIVTPR	1967	216	52	84		349	1840	
LPVPGVLLKEFTVSGNI LTI	1968	216	32	04		349	1040	
WITQCFLPVFLAQPPSG	1000	13208	23649	726		688	286	
QRR DHRQLQLSISSCLQQLS	1969	>98522.17	69	67		532	63772	
LLM	<u>1970</u>	2254	2012	0.0		1066	641	
YLAMPFATPMEAELAR RSLA	1971	3754	2813	865		1965	041	
AAPLLLARAASLSLG	1972	100	3.2	35	10470	79	79	
APLLLARAASLSLGF	1973	322	12	91	13359	59	114	
PLLLARAASLSLGFL	1974	1255	12	118	>9742.79	52	151	
SLSLGFLFLLFFWLD	1975	100000	639	11375	3710	>10955.8	66667	
LLFFWLDRSVLAKEL	1976	154	24	34	86	7.5	134	
DRSVLAKELKFVTLV	1977	20966	4410	1359	>14413.38	53	2217	
AKELKFVTLVFRHGD	1978	12309	824	1529	8563	51	24	
RSPIDTFPTDPIKES	1979	>200000	>35000	2373	>14413.38	469	28571	
FGQLTQLGMEQHYEL	1980	27217	>35000	>22750	>14413.38	543	100000	
DRTLMSAMTNLAALF	1981	2367	114	871	3927	57	26138	
MSAMTNLAALFPPEG		>200000	249	12384	7158	1072	63246	
MTNLAALFPPEGVSI	1982	141421	1310	10370	>8829.24	4606	141421	
PEGVSIWNPILLWQP	1983 1984	30861	444	7.2	4624	107	22222	
GVSIWNPILLWQPIP		10287	207	5.0	4428	492	523	
	1985							

		HL	A-DR SUP	ERTYPE				
C	SEQ ID NO.	DRB1 *1101	DRB1 *1302	DRB1 *1501	DRB3 *0101	DRB4 *0101	DRB5 *0101	DRB5 *0201
Sequence WNPILLWOPIPVHTV		19640	2259	14	>8829.24	81	100000	0201
NPILLWQPIPVHTVP	1986	599	250	4.6	>8829.24	67	25000	
PILLWQPIPVHTVPL	1987	4041	567	6.9	>8829.24	106	41491	
ILLWOPIPVHTVPLS	1988	2343	1111	65	>8829.24	712	28768	
WQPIPVHTVPLSEDQ	1989	>66666.67	2692	>45500	>8829.24	1228	>100000	
LSGLHGQDLFGIWSK	<u>1990</u>	30151	>35000	32173	>8829.24	135	81650	
YDPLYCESVHNFTLP	<u>1991</u>	30151	>35000	2136	>8829.24	6901	28768	
LPSWATEDTMTKLRE	1992	>66666.67	>35000	>45500	5973	>11134.5	343	
LISWATEDIMIKERE	1993					7		
LRELSELSLLSLYGI	1994	6958	3218	235	>14956.63	544	5185	
LSELSLLSLYGIHKQ	1995	1657	1253	45	>13046.31	79	7.3	
LSLLSLYGIHKQKEK	1996	742	>35000	58	>14956.63	772	3.4	
KSRLQGGVLVNEILN		>66666.67	318	>30333.3	>14956.63	713	>100000	
GGVLVNEILNHMKRA	1997	255	49	3 576	8124	5.8	8.7	
IPSYKKLIMYSAHDT	1998	53	2122	17	9982	12	191	
YKKLIMYSAHDTTVS	1999	208	37	15	13224	5.8	5482	
LIMYSAHDTTVSGLO	2000	>66666.67	1752	184	6828	4381	>100000	
DTTVSGLQMALDVYN	2001	>50000	3500	1042	10843	961	>200000	
ALDVYNGLLPPYASC	2002	182	>35000	1091	>14956.63	>10090.4	115470	
LDVYNGLLPPYASCH	2003	194	>35000	3035	>14956.63	7 >10918.6	25820	
and a new account	2004	5200	11447	252	>14956.63	7 >10918.6	100000	
YNGLLPPYASCHLTE	2005	5300	11667 >35000	252 >45500	>14956.63	>10918.6 7 983	>200000	
FAELVGPVIPQDWST	2006	>50000						
TVPLSEDQLLYLPFR	2007	26455	5300	>2367.33	4323	872	27221	
LTELYFEKGEYFVEM	2008	>18903.59	3157	>2367.33	124	601	6655	
GPVIPQDWSTECMTT	2009				20295	961		
QAHSLERVCHCLGKWL GHPDK	2010	2857					2500	
WTTCQSIAFPSKTSASIG SL	2011	40000		277	37450	505	400	
QKGRGYRGQHQAHSLE RVCH	2012	30151		>9100	>500000	17951	9759	
AATYNFAVLKLMGRGT KF	2013	17		239	70014	1218	18	
VATGLCFFGVALFCGC		33333			117851	193333		
GHEA FLYGALLLAEGFYTTG	2014			45			256	
AVRQ	2015						20000	
SAVPVYIYFNTWTTCQS IAF	2016			92			20000	
TLSVTWIGAAPLILS	2017	6860	642	97	6031	3506	31	
SVTWIGAAPLILSRI	2018	2196	420	147	13676	42	104	
VTWIGAAPLILSRIV	2019	1779	2339	552	>10729.61	88	147	
SQPWQVLVASRGRAV	2020	135	32	11259	>12116.81	7562	84	
GRAVCGGVLVHPQWV	2021	>50000	5456	12888	>12116.81	62	100000	
GVLVHPQWVLTAAHC	2022	263	2427	66	>10729.61	6.2	1062	
HPQWVLTAAHCIRNK	2023	785	1170	6500	1324	5518	40	
QWVLTAAHCIRNKSV	2024	2169	2062	13565	7342	3802	35	
AHCIRNKSVILLGRH	2024	93	75	88	4752	8.7	3630	
SVILLGRHSLFHPED		96	96	106	13045	4411	16116	
VILLGRHSLFHPEDT	2026	344	543	426	>12116.81	10696	100000	
GQVFQVSHSFPHPLY	2027	103	146	2172	1071	416	128	
	2028							

	SEQ	DRB1	A-DR SUPE DRB1	DRB1	DRB3	DRB4	DRB5	DRB
Sequence	ID NO.	*1101	*1302	*1501	*0101 23433	*0101 >12491.9	*0101 897	*020
VFQVSHSFPHPLYDM	2029	881	83	2396	23433	2		
PHPLYDMSLLKNRFL	2030	>50000	11667	712	>13533.63	7486	3104	
SHDLMLLRLSEPAEL	2031	4471	5.8	1099	13577	12	100000	
HDLMLLRLSEPAELT	2032	2141	2.3	662	5305	45	10541	
TDAVKVMDLPTQEPA	2033	>50000	>35000	>45500	>13533.63	747	>200000	
LHVISNDVCAQVHPQ	2034	>50000	239	22750	1887	1087	>200000	
CAQVHPQKVTKFMLC	2035	18490	2192	809	>13533.63	604	1229	
GGPLVCNGVLQGITS	2036	1828	36	30333	>6567.28	815	13417	
GPLVCNGVLQGITSW	2037	915	49	6310	11615	646	6537	
NGVLQGITSWGSEPC	2038	9724	775	258	8038	4487	11619	
RPSLYTKVVHYRKWI	2039	350	4183	717	2982	4897	13	
HSLFHPEDTGQVFQV	2040				553	11503		
PRWLCAGALVLAGGF	2040	>40000	20207	15167	13150	883	40825	
LGFLFGWFIKSSNEA		7303	10104	355	681	9285	461	
LDELKAENIKKFLYN	2042	324	597	414	548	788	150	
IKKFLYNFTQIPHLA	2043	137	27	305	477	96	658	
KFLYNFTQIPHLAGT	2044	91	221	227	10212	256	1600	
WKEFGLDSVELAHYD	2045	4935	8413	22750	829	5925	89443	
LAHYDVLLSYPNKTH	2046	380	268	82	1406	589	172	
GNEIFNTSLFEPPPP	2047	>40000	2804	>91000	>13164.82	835	>200000	
GKVFRGNKVKNAQLA	2048	894	46	3373	7591	7884	1385	
GNKVKNAOLAGAKGV	2049	>66666.67	>35000	>45500	>12462.61	1065	1218	
EYAYRRGIAEAVGLP	2050	2590	5217	>45500	8773	6325	1204	
AEAVGLPSIPVHPIG	2051	>66666.67	5456	56	>11848.34	12394	69336	
AVGLPSIPVHPIGYY	2052	33333	1191	518	>11848.34	5387	38517	
	2053	>28571.43	5729	1978	17305	13588	506	
IGYYDAQKLLEKMGG	2054			3745	>11848.34	508	1927	
TGNFSTQKVKMHIHS	2055	11856	6187	1605	17550	447	32	
TRIYNVIGTLRGAVE	2056	45	1460					
ERGVAYINADSSIEG	2057	>50000	3689	30333	6846	87	200000	
GVAYINADSSIEGNY	2058	>40000	497	7610	1420	477	66667	
DSSIEGNYTLRVDCT	2059	>50000	7.6	1202	576	1262	16824	
NYTLRVDCTPLMYSL	2060	7116	9.0	5056	25	404	66667	
CTPLMYSLVHNLTKE	2061	590	260	426	18348	58	36	
DFEVFFQRLGIASGR	2062	128	10069	10249	30745	4.2	3559	
EVFFQRLGIASGRAR	2063	31	17500	4556	>15037.59	51	7.9	
TNKFSGYPLYHSVYE	2064	33333	>35000	489	>21853.15	12466	2942	
YDPMFKYHLTVAQVR	2065	252	1014	1348	8137	553	62	
DPMFKYHLTVAQVRG	2066	69	699	230	7297	467	11	
MFKYHLTVAQVRGGM	2067	147	1615	1198	3648	1062	5.8	
KYHLTVAQVRGGMVF	2068	859	193	1222	>21853.15	3446	86	
VAQVRGGMVFELANS	2069	>50000	2802	117	>21853.15	100	64366	
RGGMVFELANSIVLP	2070	>50000	4.4	94	132	411	413	
GMVFELANSIVLPFD	2071	>50000	12	83	234	4154	903	
VFELANSIVLPFDCR	2072	11765	24	477	128	1215	10815	
ADKIYSISMKHPQEM	2073	169	4957	8273	>21853.15	3550	26726	
IYSISMKHPQEMKTY	2073	213	>35000	5025	>21853.15	5356	2588	

	SEQ	DRB1	DRB1	DRBI	DRB3	DRB4 *0101	DRB5 *0101	DRB:
Sequence PQEMKTYSVSFDSLF	ID NO.	*1101 >50000	*1302 24749	*1501 919	*0101 14564	579	100000	*020
TYSVSFDSLFSAVKN	2075	5981	5888	3223	8547	10461	61	
VLRMMNDOLMFLERA	2076	2353	130	127	98	88	85	
LRMMNDOLMFLERAF	2077	1833	1314	1411	1570	50	758	
RHVIYAPSSHNKYAG	2078	13363	8750	1291	>62814.07	5293	88	
ROIYVAAFTVOAAAE	2079	35	524	166	6808	47	143	
	2080	34	344	252	1324	50	216	
QIYVAAFTVQAAAET	2081		344 446	18200	2116	464	378	
VAAFTVQAAAETLSE	2082	2126		2713	30	3705	72993	
YISIINEDGNEIFNT	2083	>18903.59	346					
ISIINEDGNEIFNTS	2084	>18903.59	343	3006	35	6394	>37807.18	
EDFFKLERDMKINCS	2085	10433	3188	>3490.6	4036	7886	3494	
FFKLERDMKINCSGK	2086	9687	382	>3490.6	4918	98	3796	
GVILYSDPADYFAPG	2087	>18903.59	39	965	8.8	64	14168	
GAAVVHEIVRSFGTL	2088				788	89		
NSRLLQERGVAYINA	2089	12812	327	1229	3366	699	3473	
VAYINADSSIEGNYT	2090	>18903.59	2147	>3490.6	471	841	>37807.18	
DQLMFLERAFIDPLG	2091				17115	6.6		
KSNFLNCYVSGFHPSD	2092	5000					2857	
AC- NPDAENWNSQFEILED		>33333.33	>10000	>10000	1000		50000	
AA EYLILSARDVLAVVS	2093	6860		2340		2527	4154	
YKTIAYDEEARR	2094	200000		>91000	>50000		200000	
GEALSTLVVNKIRGT	2095	977	55	2314		1514	108	
PYILLVSSKVSTVKD	2096	112	7.2	22		107	32	
EAVLEDPYILLVSSK	2097	4376	>10294.12	>50837.9		>26435.7	357	
IAGLFLTTEAVVADK	2098	867	>10294.12	9 >50837.9		3 >26435.7	606	
ALSTLVVNKIRGTFK	2099	32	7.6	160		214	38	
MKHILYISFYFILVN	2100	2082					>9523.81	
KSLLSTNLPYGRTNL	2101							
HFFLFLLYILFLVKM	2102		84	21473		1064	10083	
LFLLYILFLVKMNAL	2103		129	30829		1290	32446	
ILFLVKMNALRRLPV	2104		0.13	1.4		7.6	14	
MNALRRLPVICSFLV	2105		15	36		5.7	2557	
SAFLESQSMNKIGDD	2106		52	18689		302	243	
LKELIKVGLPSFENL	2107		147	361		110	41322	
FENLVAENVKPPKVD	2108 2109		3029	>50837.9		9297	62661	
PATYGUVPVLTSLF	2110		0.83	2557		118	52	
YGIIVPVLTSLFNKV	2111		0.30	223		97	80	
LLKIWKNYMKIMNHL	2111		3.7	6.8		12	35	
MTLYQIQVMKRNQKQ			323	2429		82	22	
QKQVQMMIMIKFMGV	2113		17	363		5.3	915	
MIMIKFMGVIYIMII	2114		102	23611		145	12310	
GVIYIMIISKKMMRK	2115		38	173		157	46	
	2116							
LYYLFNOHIKKELYH	2117		327	2861		1089	606	

	SEQ	DRBI	DRB1	DRB1	DRB3	DRB4	DRB5	DRB:
Sequence	ID NO.	*1101	*1302	*1501	*0101	*0101	*0101	*0201
LDIYQKLYIKQEEQK	2119		4346	47		70	6958	
QKKYIYNLIMNTQNK	2120		53	844		87	245	
YEALIKLLPFSKRIR	2121		230	36		15	11	
ENEYATGAVRPFQAA	2122		9302	3007		10026	>10303.97	
NYELSKKAVIFTPIY	2123		410	537		136	10581	
QKILIKIPVTKNIIT	2124		332	3614		953	297	
KCLVISQVSNSDSYK	2125		236	403		81	>42553.19	
SKIMKLPKLPISNGK	2126		6460	3570		6739	>10303.97	
FIHFFTWGTMFVPKY	2127		328	2375		387	9608	
LCNFKKNIIALLIIP	2128		16	29302		99	>42553.19	
KKNIIALLIIPPKIH	2129		15	32		8.2	143	
ALLIIPPK1H1SIEL	2130		162	1823		10	7135	
SMEYKKDFLITARKP	2131		3818	4610		10448	442	
KSKFNILSSPLFNNF	2132		25	5.9		135	32	
FKKLKNHVLFLQMMN	2133		20	29		14	59	
KNHVLFLQMMNVNLQ	2134		36	224		22	>7212.41	
VLFLQMMNVNLQKQL	2135		8.6	8200		12	>7212.41	
NVNLQKQLLTNHLIN	2136		28	4448		354	>7212.41	
QKQLLTNHLINTPKI	2137		1.6	514		904	6595	
NHLINTPKIMPHHII	2138		32	560		1632	8882	
YILLKKILSSRFNQM	2139		1.01	26		340	83	
FNQMIFVSSIFISFY	2140		33	3903		1291	>12484.39	
KVSCKGSGYTFTAYQM		>200000						
H IAKVPPGPNITAEYGDK	2141	200000			>20000		200000	
WLD	2142	200000			>20000		10000	
TAEYGDKWLDAKSTW YGKPT	2143	200000			>20000		10000	
AKSTWYGKPTGAGPKD		200000			>20000		10000	
NGGA GAGPKDNGGACGYKD	2144	200000			>20000		200000	
VDKAP	2145	200000			>20000		200000	
FNGMTGCGNTPIFKDG RGCG	2146	200000			>20000		200000	
PIFKDGRGCGSCFEIKC	21.47	200000			>20000		200000	
TKP SCFEIKCTKPESCSGEA	2147	200000			>20000		200000	
VTV	2148	1010			>33333.33		200000	
AFGSMAKKGEEQNVRS AGEL	2149	1818						
TPDKLTGPFTVRYTTEG GTK	2150	200000			>25000		200000	
VRYTTEGGTKSEVEDVI	2130	200000			>25000		200000	
PEG TCVLGKLSQELHKLQ	<u>2151</u>	1398	>12589.93	2009	>263157.89	163	3986	
, ,	2152	2375	>12589.93	287	>263157.89	870	37	
KLSQELHKLQTYPRT	2153					22948	40	
LHKLQTYPRTNTGSG	2154	6091	>12589.93	157	>263157.89			
KLQTYPRTNTGSGTP	2155	8210	987	520	>263157.89	14		
CCVLGKLSQELHKLQ	2156	5243	>12589.93	570	>263157.89	346	5158	
CSNLSTCVLGKLSQE	2157	5263	7907	4538	>263157.89	11756	5709	
TSNLSTTVLGKLSQE	2158	534	9333	7697	>263157.89	13210	2529	
TTVLGKLSQELHKLQ	2159	3524	12715	525	>263157.89	241	10618	
DIAAKYKELGY	2160	>10000			>25000		200000	
ALVRQGLAKVA	216L	200000					>10000	

	cro	DRB1	A-DR SUP	ERTYPE DRB1	DRB3	DRB4	DRB5	DRB5
Sequence	SEQ ID NO.	*1101	*1302	*1501	*0101	*0101	*0101	*0201
PATLIKAIDGDTVKLMY	21/2	>6666.67			2381	,	3333	
KGQ TPETKHPKKGVEKYGP	2162	>6666.67			>25000		>4000	
EASA VEKYGPEASAFTKKMV	2163	20000			16667		34	
ENAK	2164	20000						
FTKKMVENAKKIEVEF DKGQ	2165	6667			>25000		1000	
YIYADGKMVNEALVRQ		>6666.67			>5555.56		>4000	
GLAK HEOHLRKSEAOAKKEK	2166	200000			>5555.56		11	
LNIW	2167							
QAKKEKLNIWSEDNAD SGQ	2168	200000			>5555.56		200000	
YFNNFTVSFWLRVPK	2169							
FSYFPSI	2170							
YSFFPS1	2170							
YSYFPSIR		20000					>200000	
DPNANPNVDPNANPNV	2172	>12500		>7583.33		>72500	>2898.55	
NANPNANPNANP(X4)	2173							
QKWAAVVVPS	2174							
FWQLNGEELIQDMELV ETRPAG	2175							
PEFLEQRRAAVDTYC	2176	488					200000	
STORKUSP33	2170							
DYSYLQDSDPDSFQD	2178	>66666.67	>35000	>45500			>40000	
DFSYLQDSDPDSFQD	2179		>35000	>91000			>40000	
QNILFSNAPLGPQFP	2180							
QNILLSNAPLVPQFP	2181							
DYSYLQDSDPDSFQD								
KYVKQNTLKLAT	2182 2183							
P(X)KQNTLKLAT								
EEDIEIPIOEEEY	2184	>20576.13					46083	
HQAISPRTLNSPAIF	2185	33686	1036	8106	>83333.33	130	>200000	
YTDVFSLDPTFTIETT	2186							
YAGIRRDGLLLRLVD	2187							
LFFYRKSVWSKLQSI	2188	12	121	20	5915	1933	18	
RPIVNMDYVVGARTFR	2189	222	73	43	3324	160	6.6	
REKR	2190	222	73	43	3324	100	6.6	
RPGLLGASVLGLDDI	2191	>93896.71	2056	6000	30212	22038	>88888.89	
LYFVKVDVTGAYDTI	2192	221	79	9753	16	22	4962	
FAGIRRDGLLLRLVD	2193	804	1294	28	553	1670	1355	
AKTFLRTLVRGVPEY	2194	6.3	94	829	546	472	3484	
YGAVVNLRKTVVNFP	2195	89	11236	470	51496	302	36	
GTAFVQMPAHGLFPW	2196	17	2819	1.2	769	2361	43	
WAGLLLDTRTLEVQS		20960	92	3468		862	>102040.8	
RTSIRASLTFNRGFK	2197	4807	49	497		79	2 52	
	2198						8834	
RVIKNSIRLTL	2199	1740	32	4317 2579		143 198	1039	
PVIKNSIKLRL	2200	2772	77	2319	462	198		
ATSTKKLHKEPATLIKA IDG	2201	>6666.67			462		267	

TABLE 28

	М	URINI	E CLASS I SUPERT	YPE		
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
SGPSNTPPEI	2202	10	Adenovirus	EIA		
RNPRFYNL	2203	8	Artificial sequence	Consensus		
QPQRGYENF	2204	9	Artificial sequence	Consensus		Α
SEAAYAKKI	2205	9	Artificial sequence	pool consensus		Α
AYAPAKAAI	2206	9	Artificial sequence			Poly
AYAEAKAAI	2207	9	Artificial sequence			Poly
AYANAKAAI	2208	9	Artificial sequence			Poly
AYAGAKAAI	2209	9	Artificial sequence			Poly
AYAVAKAAl	2210	9	Artificial sequence			Poly
AAAAYAAM	2211	8	Artificial sequence			
AAAAYAAAAM	2212	10	Artificial sequence			
AAAANAAAM	2213	9	Artificial sequence			
AAAAANAAAM	2214	ΙI	Artificial sequence			
NAIVFKGL	2215	8	Chicken	Ova	176	
SIINFEKL	2216	8	Chicken	Ova	257	
IFYCPIAI	2217	8	Chicken	Ova	27	
KVVRFDKL	2218	8	Chicken	Ova	55	
VYSFSLASRL	2219	10	Chicken	Ova	96	
SIINFEKL	2220	8	Chicken	Ova	257	
KVVRFDKL	2221	8	Chicken	Ova	55	
SENDRYRLL	2222	9	EBV	BZLFI	209	Α
SFYRNLLWL	2223	9	Flu	HA	142	
YEANGNLI	2224	8	Flu	HA	259	Α
MGLIYNRM	2225	8	Flu	M1	128	
MGYIYNRM	2226	8	Flu	M1	128	
MGIIYNRM	2227	8	Flu	MI	128	
MGLIFNRM	2228	8	Flu	MI	128	
MGLIYNRM	2229	8	Flu	MI	128	
RMIQNSLTI	2230	9	Flu	NP	55	
RLIQNFLTI	2231	9	Flu	NP	55	
GMRQNATEI	2232	9	Flu	NP	17	
YMRVNGKWM	2233	9	Flu	NP	97	
FYIQMATEL	2234	9	Flu	NP	39	
FYIOMCTFL	2235	9	Flu	NP	39	
AYERMANIL	2236	9	Flu	NP	218	
AYQRMCNIL	2237	9	Flu	NP	218	
AYERMCTIL	2238	9	Flu	NP	218	
ASNENMETM	2239	9	Flu	NP	366	
TYQRTRALM	2240	9	Flu	NP	147	Α
TYQKTRALV	2240	9	Flu	NP	147	Α
TYQPTRALV	2242	9	Flu	NP	147	Α
TYQFTRALV	2242	9	Flu	NP	147	Α
TYQLTRALV	2243	9	Flu	NP	147	Α
SDYEGRLI	2245	8	Flu	NP	50	
MITQFESL	2245	8	Flu	NS	31	
RTFSFQLI	2240	8	Flu	NS	114	
FSVIFDRL	2248	8	Flu	NS	134	
	2240					

	SEO ID		CLASS I SUPE			
Sequence	NO.	AA	Organism	Protein	Position	Analog
RTFSFQLI	2249	8	Flu	NS1	114	
MITQFESL	2250	8	Flu	NSI	31	
FSVIFDRL	2251	8	Flu	NS2	134	
KSSFYRNL	2252	8	FluA	HA	158	
SSLPFQNI	2253	8	FluA	HA	305	
MNIQFTAV	2254	8	FluA	HA	403	
MNYYWTLL	2255	8	FluA	HA	244	
SFYRNLLWL	2256	9	FluA	HA	160	
SSLPFQNI	2257	8	FluA	HA	305	
MNIQFTAV	2258	8	FluA	HA	403	
MNYYWTLL	2259	8	FluA	HA	244	
KSSFYRNL	2260	8	FluA	HA	158	
SIIPSGPL	2261	8	FluA	Ml	13	
LSYSAGAL	2262	8	FluA	M1	117	
LSYSAGAL	2263	8	FluA	Ml	117	
SSISFCGV	2264	8	FluA	NM	426	
TGICNQNII	2265	9	FluA	NM	46	
ITYKNSTWV	2266	9	FluA	NM	54	
FCGVNSDTV	2267	9	FluA	NM	430	
TGICNONII	2268	9	FluA	NM	46	
FCGVNSDTV	2269	9	FluA	NM	430	
ITYKNSTWV	2270	9	FluA	NM	54	
SSISFCGV	2270	8	FluA	NM	426	
IGRFYIQM	2272	8	FluA	NP	36	
MMIWHSNL	2272	8	FluA	NP	136	
ASNENMETM	2274	9	FluA	NP	366	
IGRFYIOM	2275	8	FluA	NP	36	
MMIWHSNL	2276	8	FluA	NP	136	
FFYRYGFV	2277	8	FluA	POLI	495	
KMITQRTI	2277	8	FluA	POLI	198	
RSYLIRAL	2278	8	FluA	POLI	215	
RFYRTCKL	2279	8	FluA	POLI	465	
TALANTIEV	2280	9	FluA	POLI	141	
TALANTIEV		9	FluA	POLI	141	
RSYLIRAL	2282 2283	8	FluA	POLI	215	
RFYRTCKL		8	FluA	POLI	465	
VYINTALL	2284	8	FluA	POL2	463	
VYINTALL	2285	8	FluA	POL2	463	
VYIEVLHL	2286	8	FluA	POL3	227	
VYIEVLHL	2287	8	FluA	POL3	227	
WYIPPSLRTL	2288	10	GAD	1023		
MURTAZAKDPEPTIDE	<u>2289</u>	0	GAD65		107	
S IYSTVASSL	2291	9	НА		553	
LYEKVKSQL	2291	9	HA		462	
LYQKVKSQL	2292	9	HA		462	
LYEKMKSQL	2293 2294	9	HA		462	
LYEKVFSQL	2294	9	HA		462	
LYQNVGTYV	2295 2296	9	НА		204	

		UKINI	CLASS I SUPE	KITPE		
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
MGLKFRQL	2297	8	HBV	core	122	
VSYVNTNM	2298	8	HBV	core	115	
SYVNTNMGL	2299	9	HBV	core	116	
MGLKFRQL	2300	8	HBV	core	122	
VSYVNTNM	2301	8	HBV	core	115	
SYVNTNMGL	2302	9	HBV	соге	116	
WGPSLYSI	2302	8	HBV	env	364	
ASARFSWL	2304	8	HBV	env	329	
WGPSLYSIL	2304	9	HBV	env	364	
TGPCRTCMT	2305	9	HBV	env	281	
WYWGPSLYSI	2307	10	HBV	env	362	
IPOSLDSWWTSL	2307	12	HBV	env	28	
IPQSLDSYWTSL	2308	12	HBV	env	28	Α
ASARFSWL		8	HBV	env	329	
WYWGPSLYSI	2310	10	HBV	env	362	
APOSLDSWWTSL	2311	12	HBV	env	28	
IPQALDSWWTSL	2312	12	HBV	env	28	Α
IPQSLASWWTSL	2313	12	HBV	env	28	A
IPQSLDAWWTSL	2314	12	HBV	env	28	A
-	<u>2315</u>	12	HBV	env	28	A
IPQSLDSAWTSL	<u>2316</u>	12	HBV	env	28	A
IPQSLDSWWASL	<u>2317</u>	12	HBV		28	A
IPQSLDSWWTAL	<u>2318</u>			env	28	A
EPQSLDSWWTSL	<u>2319</u>	12	HBV	env		
IPESLDSWWTSL	<u>2320</u>	12	HBV ·	env	28	A
IPQSLDEWWTSL	<u>2321</u>	12	HBV	env	28	A
IPQSLDSWWTEL	2322	12	HBV	env	28	A
RPQSLDSWWTSL	2323	12	HBV	env	28	A
IPRSLDSWWTSL	2324	12	HBV	env	28	A
IPQRLDSWWTSL	2325	12	HBV	env	28	A
IPQSRDSWWTSL	<u>2326</u>	12	HBV	env	28	A
IPQSLRSWWTSL	2327	12	HBV	env	28	Α
IPQSLDRWWTSL	2328	12	HBV	env	28	Α
IPQSLDSRWTSL	2329	12	HBV	env	28	Α
IPQSLDSWWRSL	2330	12	HBV	env	28	Α
IPQSLDSWWTRL	2331	12	HBV	env	28	Α
YPQSLDSWWTSL	2332	12	HBV	env	28	Α
IPYSLDSWWTSL	2333	12	HBV	env	28	Α
IPQYLDSWWTSL	2334	12	HBV	env	28	Α
IPQSLYSWWTSL	2335	12	HBV	env	28	Α
IPQSLDYWWTSL	2336	12	HBV	env	28	Α
IPQSLDSWYTSL	2337	12	HBV	env	28	Α
1PQSLDSWWTYL	2338	12	HBV	env	28	Α
IPGSLDSWWTSL	2339	12	HBV	env	28	Α
IPQSLDSGWTSL	2340	12	HBV	env	28	Α
IPQSLDSPWTSL	2341	12	HBV	env	28	Α
IPQSLDSWGTSL	2342	12	HBV	env	28	Α
IPQSLDSWPTSL	2343	12	HBV	env	28	Α
IPQSLDSWWTGL	2344	12	HBV	env	28	Α

		JAME	E CLASS I SUPE			
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
IPQSLDSWWTPL	2345	12	HBV	env	28	A
IPQVLDSWWTSL	2346	12	HBV	env	28	Α
IPOFLDSWWTSL	2347	12	HBV	env	28	Α
IPOPLDSWWTSL	2348	12	HBV	env	28	Α
IPOMLDSWWTSL	2349	12	HBV	env	28	Α
IPQILDSWWTSL	2350	12	HBV	env	28	Α
IPQLLDSWWTSL	2351	12	HBV	env	28	Α
IPQGLDSWWTSL	2352	12	HBV	env	28	Α
IPQTLDSWWTSL	2352	12	HBV	env	28	Α
IPOHLDSWWTSL	2354	12	HBV	env	28	A
IPOCLDSWWTSL	2354	12	HBV	env	28	A
IPONLDSWWTSL	2356	12	HBV	env	28	A
PQQLDSWWTSL		12	HBV	env	28	A
IPOWLDSWWTSL	2357	12	HBV	env	28	A
IPQDLDSWWTSL	2358	12	HBV	env	28	A
IPQKLDSWWTSL	2359	12	HBV	env	28	Ā
IPQKLDSWWTSL IPQSLVSWWTSL	2360	12	HBV	env	28	A
-	<u>2361</u>	12	HBV	env	28	Ā
IPQSLFSWWTSL IPOSLPSWWTSL	<u>2362</u>		HBV	env	28	A
•	<u>2363</u>	12	HBV		28 28	A
IPQSLMSWWTSL	<u>2364</u>	12		env	28	A
IPQSLISWWTSL	2365	12	HBV	env		
IPQSLLSWWTSL	<u>2366</u>	12	HBV	env	28	A
IPQSLGSWWTSL	<u>2367</u>	12	HBV	env	28	A
IPQSLSSWWTSL	2368	12	HBV	env	28	A
IPQSLTSWWTSL	<u>2369</u>	12	HBV	env	28	A
IPQSLHSWWTSL	2370	12	HBV	env	28	A
IPQSLCSWWTSL	2371	12	HBV	env	28	Α
IPQSLNSWWTSL	2372	12	HBV	env	28	Α
IPQSLQSWWTSL	2373	12	HBV	env	28	Α
IPQSLWSWWTSL	2374	12	HBV	env	28	Α
IPQSLKSWWTSL	2375	12	HBV	env	28	Α
IPSLDSWWTSL	2376	11	HBV	env	28	Α
IPQSLDSWTSL	2377	11	HBV	env	28	Α
IPQSLDSWWTL	2378	11	HBV	env	28	Α
IPQALASWWTSL	2379	12	HBV	env	28	Α
IPQSLDSWWTSM	2380	12	HBV	env	28	Α
IPQSLDSWWTSF	2381	12	HBV	env	28	Α
KTPSFPNI	2382	8	HBV	pol	75	
HAVEFHNL	2383	8	HBV	pol	289	
VSAAFYHL	2384	8	HBV	pol	419	
VIGCYGSL	2385	8	HBV	pol	588	
KQYLNLYPV	2386	9	HBV	pol	668	
CYGSLPQEHI	2387	10	HBV	pol	591	
VSAAFYHL	2388	8	HBV	pol	419	
HAVEFHNL	2389	8	HBV	pol	289	
VIGCYGSL	2390	8	HBV	pol	588	
KTPSFPNI	2391	8	HBV	pol .	75	
RPQSLDSWWTSL	2392	12	HBVs	env	28	Α

		IURINI	E CLASS I SUPE	RIYPE		
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
IPQRLDSWWTSL	2393	12	HBVs	env	28	Α
IPQSLRSWWTSL	2394	12	HBVs	env	28	Α
IPQSLDRWWTSL	2395	12	HBVs	env	28	Α
IPOSLDSRWTSL	2396	12	HBVs	env	28	Α
IPQSLDSWWRSL	2397	12	HBVs	env	28	Α
IPQSLDSWWTRL	2398	12	HBVs	env	28	Α
IPOELDSWWTSL	2399	12	HBVs	env	28	Α
IPQSLYSWWTSL	2400	12	HBVs	env	28	Α
IPQSLDSWETSL	2401	12	HBVs	env	28	Α
IPQSLDSWWESL	2402	12	HBVs	env	28	Α
VESENKVV	2403	8	HCV	Entire	2253	
AGPYRAFVTI	2404	10	HIV	env	18	Α
RAPYRAFVTI	2404	10	HIV	env	18	Α
RGPYRAFVTA	2405	10	HIV	env	18	Α
KGPYRAFVTI	2406 2407	10	HIV	env	18	A
RGPYRAFVTK	2407	10	HIV	env	18	A
RGPGRAFVTI	2408 2409	10	HIV	env	18	
RGPGRYFVTI		10	HIV	env	18	Α
RGPGRAYVTI	2410	10	HIV	env	18	A
RGPGRAFYTI	2411	10	HIV	env	18	A
VESMNKEL	2412	8	HIV	POL	903	
TDSQYALGI	<u>2413</u> 2414	9	HIV	POL	689	
RGAYRAFVTI		10	HIV		18	Α
RGPARAFVTI	2415	01	HIV		18	A
RGPYRAAVTI	2416	10	HIV		18	A
RGPYRAFATI	2417	10	HIV		18	A
RGPYRAFVAI	2418	10	HIV		18	A
RGKYRAFVTI	<u>2419</u>	01	HIV		18	A
RGPFRAFVTI	2420	10	HIV		18	A
RGPYKAFVTI	2421	10	HIV		18	A
RGPYRKFVTI	2422	10	HIV		18	A
RGPYRAYVTI	2423	01	HIV		18	A
RGPYRAFKTI	2424	10	HIV		18	A
RGPYRAFVKI	2425	10	HIV		18	A
NEILIRCII	2426	9	HPV	E6	97	^
QEKKRHVDL	2427	9	HPV	E6	113	
LFVVYRDSI	2428	9	HPV	E6	52	
FYSRIRELRF	2429	10	HPV	E6	71	Α
SSIEFARL	2430	8	HSV	20	498	
KVPRNQDWL	2431	9	Human	gp100	1,70	
VYDFYVWM	2432	8	Human	TRP2		Α
KNKFFSYL	2433	8	Human	Tyrosinase	131	
LAVLYCLL	2434	8	Human	Tyrosinase	3	
YMVPFIPL	2435	8	Human	Tyrosinase	425	
GQMNNGSTPM	2436	10	Human	Tyrosinase	157	
IVTMFEAL	2437	8	LCMV	GP	4	
ISHNFCNL	2438	8	LCMV	GP	118	
GVYQFKSV	2439 2440	8	LCMV	GP	70	
GVYQFKSV	2440	٥	LCIVI V	GF	70	

		URINI	E CLASS I SUPER	RTYPE		
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
HYISMGTSGL	2441	10	LCMV	GP	99	,,,,,,,,,,
SGVENPGGYCL	2442	11	LCMV	GP	276	
KAVYNFATM	2442 2443	9	LCMV	GP	33	
CMANNSHHYI		10	LCMV	GP	92	Α
CSANNSHHYM	2444	10	LCMV	GP	92	A
SMVENPGGYCL	2445	11	LCMV	GP	276	A
SGVENPGGYCM	2446	11	LCMV	GP	276	A
KAVYNFATM	2447	9	LCMV	GP	33	
KAVYNAATM	2448	9	LCMV	GP GP	33	Α
	<u>2449</u>	9	LCMV	GP	33	A
KAVANFATM	<u>2450</u>	9	LCMV	GP GP	33	A
KAVYNYATM	<u>2451</u>	-				
KAVYNFAAM	<u>2452</u>	9	LCMV	GP	33	Α
YTVKYPNL	<u>2453</u>	8	LCMV	NP	205	
FQPQNGQFI	<u>2454</u>	9	LCMV	NP	396	
VGLSYSQTM	<u>2455</u>	9	LCMV	NP	356	
FQPQNGQFI	2456	9	LCMV	NP	396	
FQPQNGQFIHFY	2457	12	LCMV	NP	396	
RPQASGVYM	2458	9	LCMV	NP	118	
RPQASQVYM	2459	9	LCMV	NP	118	Α
YTYKYPNL	2460	8	LCMV	NP	205	Α
RPQASGVYM	2461	9	LCMV	NP	118	Α
RPQASGVAM	2462	9	LCMV	NP	118	Α
RPQGSGVYM	2463	9	LCMV	NP	118	Α
RPNASGVYM	2464	9	LCMV	NP	118	Α
KAVYNFATCGI	2465	11	LCMV			
KAVYNFATB	2466	9	LCMV			
VYAKECTGL	2467	9	Lysteria	listeriolysin	479	
YPHFMPTNL	2468	9	MCMV		168	
YPHYMPTNL	2469	9	MCMV		168	Α
HETTYNSI	2470	8	Mouse	beta actin	275	Α
YEDTGKTI	2410	8	Mouse	p40 phox	245	
	2471			RNA		
LGYDYSYL	2472	8	Mouse	Tyrosinase	445	
SSMHNALHI	2473	9	Mouse	Tyrosinase	360	
ANFSFRNTL	2474	9	Mouse	Tyrosinase	336	
SYLTLAKHT	2475	9	Mouse	Tyrosinase	136	
HYYVSRDTL	2476	9	Mouse	Tyrosinase	180	
YYVSRDTLL	2477	9	Mouse	Tyrosinase	181	
SFFSSWQII	2478	9	Mouse	Tyrosinase	267	
SYMVPFIPL	2479	9	Mouse	Tyrosinase	424	
PYLEQASRI	2480	9	Mouse	Tyrosinase	466	
SYLTLAKHTI	2481	10	Mouse	Tyrosinase	136	
HYYVSRDTLL	2482	10	Mouse	Tyrosinase	180	
SQVMNLHNL	2483	9	Mouse	TYRP2	363	
YENDIEKKI	2484	9	P. falciparum	CSP	375	
NEEPSDKHI	2485	9	P. falciparum	CSPZ	347	
EEKHEKKHV	2486	9	P. falciparum	LSA1	52	
SYVPSAEQIL	2480 2487	10	P. yoelii	CSP	280	
	2707	10	Unknown	HLA-A24	170	

	M	URINI	CLASS I SUPE	RTYPE		
Sequence	SEQ ID NO.	AA	Organism	Protein	Position	Analog
RYLKNGKETL	2489	10	Unknown	HLA-Cw3	170	
IYTQNRRAL	2490	9	Unknown	P815	12	
VYDFFVWM	2491	8	Unknown	TRP2	181	Α
SVYDFFVWL	2492	9	Unknown	TRP2	180	
SVYDFYVWM	2493	9	Unknown	TRP2	180	Α
ASNENMDAM	2494	9	unknown			
FAPGYNPAL	2495	9	unknown			
SIQFFGERAL	2496	10	unknown			
SIQFFGEL	2497	8	unknown			
RGYVYQGL	2498	8	VSV	NP	52	
RGPRLNTL	2499	8				
HMWNFIGV	2500	8				
GGAYRLIVF	2501	9				
KYLVTRHADV	2502	19				
FSPRRNGYL	2503	9				
SHYAFSPM	2177	8				
FQPQNGQFI	2290	9				

TABLE 29

	SEQ	MURIN	C CLASS	ISUPER	LIFE		
	ID						
Sequence	NO.	Dd	Kb	Kd	Db	Ld	Kk
SGPSNTPPEI	2202	18500	>31000	>10000	8.1		
RNPRFYNL	2203		7.9		>44000		
QPQRGYENF	2204					319	
SEAAYAKKI	2205						3.9
AYAPAKAAI	2206			3.5			
AYAEAKAAI	2207			50			
AYANAKAAI	2208			60			
AYAGAKAAI	2209			48			
AYAVAKAAI	2210			42			
AAAAYAAM	2211		375		>44000		
AAAAYAAAAM	2212		228		>44000		
AAAANAAAM	2213		10960		23		
AAAAANAAA			31000		257		
M	2214		484				
NAIVFKGL	2215						
SIINFEKL	<u>2216</u>		3.7				
IFYCPIAI	2217		195				
KVVRFDKL	2218		92	202			
VYSFSLASRL	2219			303	20500		
SIINFEKL	2220	>37000	1.5	>10000	30508		
KVVRFDKL	2221		37				
SENDRYRLL	2222						13
SFYRNLLWL	2223			>10000	304		
YEANGNLI	2224						0.65
MGLIYNRM	2225		16				
MGYIYNRM	2226		2.3				
MGIIYNRM	2227		14				
MGLIFNRM	2228		21				
MGLIYNRM	2229		9.9				
RMIQNSLTI	2230				4.6		
RLIQNFLTI	2231				40		
GMRQNATEI	2232				81		
YMRVNGKWM	2233				50		
FYIQMATEL	2234			0.31			
FYIQMCTFL	2235			1.1			
AYERMANIL	2236			233			
AYQRMCNIL	2237			2.7			
AYERMCTIL	2238			4. I			
ASNENMETM	2239	>37000	>31000	>10000	33		
TYQRTRALM	2240			69			
TYQKTRALV	2241			44			
TYQPTRALV	2242			17			
TYQFTRALV	2243			371			
TYQLTRALV	2244			110			
SDYEGRLI	2245						0.60
MITQFESL	2246		64				
RTFSFQLI	2247		26				

	1	MURINE	CLASS	ISUPE	RTYPE		
	SEQ						
Camanaa	<u>ID</u> NO.	Dd	Кb	Kd	Db	Ld	Kk
Sequence FSVIFDRL		Du	201	Ku		Lu	- IXK
RTFSFQLI	2248		27				
MITOFESL	2249		42				
FSVIFDRL	2250		115				
KSSFYRNL	2251		209				
	2252		53				
SSLPFQNI	2253						
MNIQFTAV	2254		131				
MNYYWTLL	2255		169		40		
SFYRNLLWL	2256				46		
SSLPFQNI	2257		9.5				
MNIQFTAV	2258		26				
MNYYWTLL	2259		56				
KSSFYRNL	2260		117				
SIIPSGPL	2261		393				
LSYSAGAL	2262		60				
LSYSAGAL	2263		31				
SSISFCGV	2264		29				
TGICNQNII	2265				13		
ITYKNSTWV	2266				409		
FCGVNSDTV	2267				206		
TGICNQNII	2268				21		
FCGVNSDTV	2269				166		
ITYKNSTWV	2270				276		
SSISFCGV	2271		2.3				
IGRFYIQM	2272		42				
MMIWHSNL	2273		238				
ASNENMETM	2274				41		
IGRFYIQM	2275		24				
MMIWHSNL	2276		287				
FFYRYGFV			350				
KMITQRTI	2277		300				
RSYLIRAL	2278		103				
RFYRTCKL	2279		117				
TALANTIEV	2280				16		
TALANTIEV	2281				3.7		
RSYLIRAL	2282		78		5.7		
RFYRTCKL	2283		47				
VYINTALL	2284		65				
VYINTALL	2285		14				
	2286						
VYIEVLHL	2287		75				
VYIEVLHL	2288		21	06			
WYIPPSLRTL	2289			96			
MURTAZAKDPE PTIDES				0.96 4.1			
IYSTVASSL	2291						
LYEKVKSQL	2292			2.2			
LYQKVKSQL	2293			2.8			
LYEKMKSQL	2294			1.6			

		MURIN	E CLASS	ISUPER	RTYPE		
	SEQ ID						
Sequence	NO.	Dd	Kb	Kd	Db	Ld	Kk
LYEKVFSQL	2295			7.4			
LYQNVGTYV	2296			6.9			
MGLKFRQL	2297		7.4				
VSYVNTNM	2298		60				
SYVNTNMGL	2299			19			
MGLKFRQL	2300		6.3				
VSYVNTNM	2301		33				
SYVNTNMGL	2302			12			
WGPSLYSI	2303	17					
ASARFSWL	2304		323				
WGPSLYSIL	2305	6.6					
TGPCRTCMT	2306	108					
WYWGPSLYSI	2307			8.3			
IPQSLDSWWTS	2308					2.2	
IPQSLDSYWTSL	2309					2.7	
ASARFSWL	2310		49				
WYWGPSLYSI	2311			16			
APQSLDSWWTS	2311					15	
L	2312						
IPQALDSWWTS L	2313					6.1	
IPQSLASWWTS	2313					4.2	
L	2314						
IPQSLDAWWTS L	2315					4.0	
IPQSLDSAWTSL	2316					13	
IPQSLDSWWAS	22.0					0.34	
L	2317						
IPQSLDSWWTA L	2318					134	
EPQSLDSWWTS	2310					86	
L	2319						
IPESLDSWWTSL	2320					13	
IPQSLDEWWTS L	2321					1.9	
IPQSLDSWWTE	2021					3.0	
L	2322						
RPQSLDSWWTS L	2323					60	
IPRSLDSWWTS	2323					160	
L	2324						
IPQRLDSWWTS L	2325					23	
IPQSRDSWWTS	and the					21	
L	2326						
IPQSLRSWWTS	2327					12	
IPQSLDRWWTS	****					5.0	
L	2328					47	
IPQSLDSRWTSL	<u>2329</u>					47	
IPQSLDSWWRS L	2330					485	
IPQSLDSWWTR	222V					196	
L VBOST DSWWTE	2331					91	
YPQSLDSWWTS L	2332					71	

	SEQ	TORINI	CLASS	ISUPER	E		
	ID						
equence	NO.	Dd	Kb	Kd	Db	Ld	Kk
PYSLDSWWTS						0.78	
PQYLDSWWTS	<u>2333</u>					92	
	2334						
PQSLYSWWTS						4.7	
, DOOL BANKENER	2335					1.6	
PQSLDYWWTS	2336					1.0	
PQSLDSWYTSL	2337					17	
POSLDSWWTY	2231					0.89	
,	2338						
PGSLDSWWTS						24	
, PQSLDSGWTSL	2339					70	
-	2340					19	
PQSLDSPWTSL	<u>2341</u>					138	
PQSLDSWGTSL	2342					60	
POSLDSWPTSL	2343					2.5	
PQSLDSWWTG	2344					2.3	
PQSLDSWWTP	25.22					1.2	
	2345						
PQVLDSWWTS						5.1	
, PQFLDSWWTS	<u>2346</u>					4.3	
QI LDSW W 15	2347					4.5	
PQPLDSWWTS						6.3	
DOLLI DOLLUTO	<u>2348</u>					4.1	
PQMLDSWWTS	2349					4.1	
PQILDSWWTSL	2350					12	
PQLLDSWWTS	2000					0.25	
	2351						
PQGLDSWWTS	2252					2.7	
, PQTLDSWWTS	2352					7.7	
,	2353						
PQHLDSWWTS						39	
PQCLDSWWTS	<u>2354</u>					25	
, CCLDSWW13	2355					23	
PQNLDSWWTS						12	
, DOOL DOUBLETS	2356					1.7	
PQQLDSWWTS	2357					1.7	
PQWLDSWWTS	4001					3.7	
	2358						
PQDLDSWWTS	2250					22	
, PQKLDSWWTS	2359					9.3	
, QREEDON III TO	2360						
PQSLVSWWTS						11	
POSI FSWWTSI	2361					11	
PQSLFSWWTSL	2362					16	
PQSLPSWWTSL	2363					0.95	
PQSLMSWWTS	2364					0.93	
						17	
	2365					17	
PQSLISWWTSL PQSLLSWWTSL	2365 2366					0.84	

	SEQ ID		E CLASS				
Sequence	NO.	Dd	Kb	Kd	Db	Ld	Kk
,						0.40	
PQSLSSWWTSL	2368					0.49	
PQSLTSWWTSL	2369					1.7	
PQSLHSWWTS	2370					1.5	
PQSLCSWWTS	23/0					1.1	
	2371						
PQSLNSWWTS	2272					1.5	
PQSLQSWWTS	2372					0.81	
	2373						
PQSLWSWWTS						2.4	
PQSLKSWWTS	2374					1.1	
L	2375					•••	
PSLDSWWTSL	2376					119	
PQSLDSWTSL	2377					0.22	
PQSLDSWWTL	2378					1.3	
PQALASWWTS						26	
DOG! DOUBLETS	2379					0.80	
PQSLDSWWTS M	2380					0.80	
PQSLDSWWTS						1.9	
	2381						
KTPSFPNI	2382		270				
HAVEFHNL	2383		49				
/SAAFYHL	2384		7.0				
VIGCYGSL	2385		157				
KQYLNLYPV	2386				3.4		
CYGSLPQEHI	2387			303			
VSAAFYHL	2388		5.2				
HAVEFHNL	2389		158				
VIGCYGSL	2390		63				
CTPSFPNI	2391		155				
RPQSLDSWWTS	2202					144	
PQRLDSWWTS	2392					34	
L	2393						
PQSLRSWWTS	2204					11	
L POSLDRWWTS	2394					2.0	
L	2395						
PQSLDSRWTSL	2396					2.6	
PQSLDSWWRS						335	
POSLDSWWTR	2397					27	
rQ3LD3WWIK	2398						
PQELDSWWTS						18	
DOG! VEWNITE	2399					8.3	
PQSLYSWWTS	2400					0.3	
PQSLDSWETSL	2401					5.3	
PQSLDSWWES						394	
ALCONINATA	2402						349
VESENKVV	2403	5.0					545
AGPYRAFVTI	2404	5.0					

		MURIN	E CLASS	I SUPER	TYPE		
	SEQ						
Cognonac	<u>ID</u> NO.	Dd	Кb	Kd	Db	Ld	Kk
Sequence RAPYRAFVTI		176	- KU	Ku	DU		
RGPYRAFVTA	2405	126					
KGPYRAFVTI	2406	5.8					
RGPYRAFVTK	2407	91					
RGPGRAFVTI	2408	9.7	31000	>10000	22000		
RGPGRYFVTI	2409	2.7	31000	-10000	22000		
RGPGRAYVTI	2410	14					
	2411						
RGPGRAFYTI	2412	7.2					114
VESMNKEL	<u>2413</u>						179
TDSQYALGI	2414						179
RGAYRAFVTI	2415	3.4					
RGPARAFVTI	2416	1.04					
RGPYRAAVTI	2417	2.0					
RGPYRAFATI	2418	2.1					
RGPYRAFVAI	2419	1.3					
RGKYRAFVTI	2420	67					
RGPFRAFVTI	2421	0.78					
RGPYKAFVTI	2422	13					
RGPYRKFVTI	2423	3.6					
RGPYRAYVTI	2424	2.1					
RGPYRAFKTI	2425	2.3					
RGPYRAFVKI	2426	3.9					
NEILIRCII	2427						12
QEKKRHVDL	2428						256
LFVVYRDSI	2429			453			
FYSRIRELRF	2430			447			
SSIEFARL	2431		1.8	>10000			
KVPRNQDWL	2432				38		
VYDFYVWM	2433		145				
KNKFFSYL	2434		57				
LAVLYCLL	2435		72				
YMVPFIPL	2436		70				
GQMNNGSTPM	2437				242		
IVTMFEAL	2438		82				
ISHNFCNL	2439		411				
GVYQFKSV	2440		11				
HYISMGTSGL	2441			83			
SGVENPGGYCL	2442		>31000		60		
KAVYNFATM	2442				3.3		
CMANNSHHYI					220		
CSANNSHHYM	2444				42		
SMVENPGGYCL	2445				154		
SGVENPGGYCM	2446				128		
KAVYNFATM	2447				1.5	>27000	
KAVYNAATM	2448				2.0	>27000	
KAVANFATM	2449				1.2	27000	
KAVYNYATM	2450				2.1	>27000	
KAVYNFAAM	2451				4.4	27000	
	2452						

		MURINE	CLASS	ISUPER	RTYPE		
_	SEQ ID						***
Sequence	NO.	Dd	<u>Кь</u> 204	Kd	Db	Ld	Kk
YTVKYPNL	2453		204		6.9		
FQPQNGQFI	<u>2454</u>				6.9		
VGLSYSQTM	<u>2455</u>		71				
FQPQNGQFI	2456		>31000		4.9		
FQPQNGQFIHFY	2457		15500		280		
RPQASGVYM	2458		>31000		>44000	0.99	
RPQASQVYM	2459					3.8	
YTYKYPNL	2460		1.8				
RPQASGVYM	2461					3.0	
RPQASGVAM	2462					12	
RPQGSGVYM	2463					39	
RPNASGVYM	2464					19	
KAVYNFATCGI	2465				29		
KAVYNFATB	2466				7.9		
VYAKECTGL	2467			129			
YPHFMPTNL	2468					7.5	
YPHYMPTNL	2469					9.5	
HETTYNSI	2470						1.8
YEDTGKTI	2471						0.86
LGYDYSYL	2472		3.4				
SSMHNALHI	2473				7.6		
ANFSFRNTL	2474		6.0				
SYLTLAKHT	2475			188			
HYYVSRDTL	2475			43			
YYVSRDTLL	2477			99			
SFFSSWQII				16			
SYMVPFIPL	2478			144			
PYLEQASRI	2479			173			
SYLTLAKHTI	2480			4.4			
HYYVSRDTLL	2481			167			
SOVMNLHNL	2482			107	2.3		
YENDIEKKI	2483				2.3		3.8
NEEPSDKHI	<u>2484</u>						40
EEKHEKKHV	<u>2485</u>						284
	2486			280			204
SYVPSAEQIL	<u>2487</u>			80			
RYLENGKETL	2488						
RYLKNGKETL	<u>2489</u>			217			
IYTQNRRAL	<u>2490</u>			144			
VYDFFVWM	2491		464				
SVYDFFVWL	2492		1.0				
SVYDFYVWM	2493		1.2		3365		
ASNENMDAM	2494				28		
FAPGYNPAL	2495		2.0				
SIQFFGERAL	2496		21		>44000		
SIQFFGEL	2497		16		>44000		
RGYVYQGL	2498	>37000	2.1	>10000	>44000		
RGPRLNTL	2499	186					
HMWNFIGV	2500		202				

MURINE CLASS I SUPERTYPE							
Sequence	SEQ ID NO.	Dd	Kb	Kd	Db	Ld	Kk
GGAYRLIVF	2501	3.5					-
KYLVTRHADV	2502			33			
FSPRRNGYL	2503	2.7					
SHYAFSPM	2177		250		>88000		
FQPQNGQFI	2290		9513		17		

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VIII-4-1	Declaration: Inventorship (only for the purposes of the designation of the United States of America)	
	Declaration of Inventorship (Rules 4.17(iv) and 51bis.1(a)(iv)) for the	I hereby declare that I believe I am the
	purposes of the designation of the United States of America:	original, first and sole (if only one inventor is listed below) or joint (if
		more than one inventor is listed below)
		inventor of the subject matter which is
		claimed and for which a patent is
		sought.
		This declaration is directed to the
		international application of which it
		forms a part (if filing declaration with
		application).
		I hereby declare that my residence,
		mailing address, and citizenship are as
		stated next to my name.
		I hereby state that I have reviewed and
	•	understand the contents of the
		above-identified international
		application, including the claims of
		said application. I have identified in
		the request of said application, in
		compliance with PCT Rule 4.10, any clair
		to foreign priority, and I have
		identified below, under the heading
		"Prior Applications," by application
		number, country or Member of the World
		Trade Organization, day, month and year
ı	•	of filing, any application for a patent
		or inventor's certificate filed in a
		country other than the United States of
- 1		America, including any PCT international
		application designating at least one
		country other than the United States of
		America, having a filing date before
		that of the application on which foreign
		priority is claimed.
III-4-1	Prior applications:	60/416,207, US, 03 October 2002
		(03.10.2002)
		60/417,269, US, 08 October 2002
		(08.10.2002)

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VIII-4-1 -2-1	Name:	SOUTHWOOD, Scott
VIII-4-1 -2-2	Residence: (city and either US State, If applicable, or country)	Santee, California
VIII-4-1 -2-3	Mailing address:	10679 Strathmore Drive
VIII-4-1 -2-4	Citizenship:	US
VIII-4-1 -2-5	Inventor's Signature: (if not contained in the request, or if declaration is corrected or added under Rule 26ter after the filing of the International application. The signature must be that of the inventor, not that of the agent) Date:	1 car South 5
-2-6	(of signature which is not contained in the request, or of the declaration that is corrected or added under Rule 26ter after the filing of the international application)	"/5/05
VIII-4-1 -3-1	Name:	SETTE, Alessandro
VIII-4-1 -3-2	Residence: (city and either US State, if applicable, or country)	La Jolla, California
VIII-4-1 -3-3	Mailing address:	5551 Linda Rosa Avenue
VIII-4-1 -3-4		IT / () (O /
VIII-4-1 -3-5	Inventor's Signature: (if not contained in the request, or if declaration is corrected or added under Rule 26ter after the filing of the international application. The signature must be that of the inventor, not that of the agent)	Alder J. J.
VIII-4-1.	Date:	11/17/83
-3-0	(of signature which is not contained in the request, or of the declaration that is	
-	corrected or added under Rule 26ter after the filing of the international application)	